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Effects of inter-varietal diversity, biotic stresses and environmental productivity on grain yield of spring barley variety mixtures

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Abstract Varietal seed mixtures tend to increase and stabilize crop yields, yet their application is sparse. Large-scale cultivation of variety mixtures may require a better understanding of how inter-varietal interactions and their interaction with the environment may influence the grain yield of variety mixtures relative to their component varieties. For this purpose, six variety mixtures of spring barley and 14 component varieties were grown in each of 17 trial environments. A total of 28 observed and a priori plant characteristics, including grain yield, disease severity and weed competitiveness, were derived for each component

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variety in each trial. The relationship between intervarietal diversity of each characteristic and the mixing effect on grain yield was analysed. Additionally, various types of yield stability were estimated and compared among mixtures and component varieties. One mixture out-yielded all of its component varieties in almost half of the trial environments. Inter-varietal diversity in grain yield potential correlated significantly with mixing effect, as did straw length diversity when weighted with weed pressure. The grain yields of most mixtures were more stable across environments than their component varieties when accounting also for the general response to environmental productivity. Hence, most mixtures adapted slightly better to environmental productivity and were less sensitive to environmental stress than their component varieties. We conclude that the efficacy of variety mixtures may be enhanced by mixing relatively high-yielding varieties differing in responsiveness to environmental productivity.

Keywords Compensation · Complementarity · Disease severity · Environmental response · Weed infestation · Yield stability

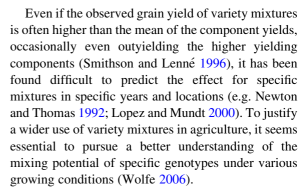
Introduction

Growth conditions of plants vary substantially between locations and years, and it is well-known that crop varieties of cereals may differ widely in their



response to these conditions. Such genotype-environment interactions (e.g. Finlay and Wilkinson 1963; Langer et al. 1979; Piepho 1998) are important contributors to unpredictability when breeders select genetic material and when farmers select varieties to be grown at a specific location (e.g. Allard and Bradshaw 1964; Ceccarelli 1996; Østergård et al. 2005; Wolfe et al. 2008). Yield stability is a key target in farm management, regional variety testing trials as well as crop breeding programs and has been defined and measured in a lot of different ways addressing different questions (for a review, see Robert 2002).

Seed mixtures of varieties with differing characteristics has been demonstrated as a potential means of increasing as well as stabilizing crop yield over environments (for reviews see Smithson and Lenné 1996; Finckh et al. 2000; Kiær et al. 2009). Specifically two types of interaction among the cooccurring varieties have been suggested and demonstrated to derive from such inter-varietal diversity within the crop stand, i.e. compensation and complementarity. First, varieties that perform well in a given environment may compensate the sub-optimal growth of others (e.g. Stützel and Aufhammer 1990), whether this results from inter-plant competition or environmental mismatch. Second, several types of varietal complementarity may positively affect crop yields: (a) complementarity in the strategy of varieties for utilizing natural resources (i.e. niche differentiation) may result in higher land use efficiency and generally better competitiveness against weeds, for example due to differences in height (e.g. Sage 1971; Kaut et al. 2009); (b) varietal complementarity in susceptibility to abiotic stress may result in generally higher and more stable grain yields of mixtures across growing environments (Smithson and Lenné 1996), e.g. varietal complementarity in the tendency to lodge under adverse weather conditions may allow more sturdy varieties to support the erect growth of others; and (c) varietal complementarity in resistance genes towards specific diseases has been shown to confer higher level of resistance of variety mixtures under a range of circumstances (Finckh et al. 2000); so far, the latter has been the primary reason for growing variety mixtures on commercial scale. Variety mixtures are thus expected to minimise the risk of reduced yield under stress conditions and may thus contribute to yield stability across growth environments.



The aim of this study was to investigate why some spring barley variety mixtures perform better than others relative to their component varieties. Therefore, we first assessed the yield potential and mixing effect of six variety mixtures over a broad range of environments (years, locations and crop management types). Second, we analysed the importance of diversity in various characteristics of the 14 component varieties for mixing success under the considered range of growing conditions, hypothesizing that larger mixing effects can be seen at higher levels of inter-varietal diversity. This hypothesis was put forward in a recent meta-analysis of a large number of mixtures and trials, which found that the characteristic most correlated with mixing effect was the diversity of component variety yields (Kiær et al. 2009). Third, we compared yield stability patterns of mixtures relative to pure stands of component varieties, using three different measures of stability. Ultimately, the results from the different analyses were combined to generate three hypotheses on how potential inter-varietal interactions may contribute to mixture yields and yield stability.

Materials and methods

Field trials

Field trials were conducted in the years 2002–2005 at four Danish locations: the three research stations Flakkebjerg (sandy loam), Foulum (loamy sand) and St. Jyndevad (coarse sand) and a certified organic farm at Dalmose (sandy loam). Trials represented one of three different low-input crop management strategies (Table 1). Herbicides and mineral fertilizers were applied to 'conventional' trials, whereas 'organic' trials comprised a variation of low input systems with different history of crop rotation and management.



Table 1 Trial environments of the study

Environment ^a	No. genotypic entities	Average grain yield (hkg ha ⁻¹
Fou05_u	35	27.4
Jyn04_u	48	35.3
Dal05_o	43	40.4
Fla04_o	48	42.0
Jyn04_o	48	47.5
Fou05_o	35	50.3
Fou04_u	48	50.6
Fla02_o	123	51.0
Jyn03_o	132	52.0
Fou02_c ^b	119	52.8
Fla03_o	132	54.9
Fou03_c	132	54.9
Fou03_o	132	55.1
Fla02_c ^b	119	56.3
Fou02_o	123	56.5
Fou04_o	48	58.5
Fla03_c	132	63.8

^a The environment coding is 3 letters for the locations Flakkebjerg, Foulum, Jyndevad and Dalmose (all in Denmark), 2 digits for year (02 denoting 2002, etc.), 1 letter for management (*u* undersown 'organic', *ο* 'organic', *c* 'conventional', see text)

Hence, the 'organic' systems comprised trials with or without undersown grass clover mixtures, the former with no nutrients added and the latter with a reduced rate of manure and a weed harrowing strategy including one pre-emergence weed harrowing and 1-3 rounds of postemergence weed harrowing to keep weed pressures manageable (see Hansen et al. 2008). Fungicides were not applied in any trial. A total of 17 combinations of year, location and management were included in this study (Table 1), hereafter designated as environments. Field plots were rectangular strips of width 1.5 m, with an area varying among trials from 12 to 20 m². Between 35 and 132 varieties and variety mixtures of spring barley (hereafter designated genotypic entities) were grown in each environment (Table 1). Each trial was laid out in an incomplete block design (α-design, Patterson et al. 1978) with two or three replicates. Each trial was analysed with a mixed effects model,

$$Y_{ijkl} = \mu_{il} + C_{kl} + D_{jkl} + E_{ijkl}, \tag{1}$$

where Y_{ijkl} is the observed grain yield of the *i*th of ν genotypic entities in the *j*th of b blocks in the kth of

r replicates in environment l, μ_{il} is the expected mean yield of genotypic entity i in environment l, while C_{kl} , D_{jkl} and E_{ijkl} are random effects of replicate, block within replicate, and residual error, each assumed to be iid normally distributed with mean value 0 and variances σ_C^2 , σ_D^2 and σ_F^2 , respectively.

Varieties and mixtures studied

Each of the six considered variety mixtures were composed of three component varieties from a set of 14 (Table 2). The varieties and mixture combinations were selected to study interactions between component varieties and the influence of variation in varietal characteristics for mixture performance, specifically with respect to height and weed competitiveness. For this reason, mixture components were chosen to have generally high disease resistance and larger than recommended inter-varietal differences in straw length. Further, some mixtures combined malting and fodder varieties. It is important to note that the mixtures were not designed to maximise mixing effects or to study the control of disease resistance. In order to ensure acceptable agronomic performance all mixtures complied with the official Danish certification requirements for mixture components concerning relative yield, disease resistance, and date of ripening (see Østergård and Jensen 2005). One older variety (Culma) was not grown in two of the environments, and the effect of mixing could not be estimated in these environments for the mixture (Mix2) to which it contributed.

Analysis of mixing effect

In order to level out differences in productivity among trial environments and the resulting variation in errors, which were observed to affect comparisons of absolute values of yield in mixture and pure stand plots, a relative measure of mixing effect was calculated for each mixture in each environment by a relative effect measure as

$$MErel_{ml} = \frac{\hat{\mu}_l^m - \hat{\mu}_{ml}^c}{\hat{\mu}_{ml}^c},$$
 (2)

where $\hat{\mu}_l^m$ is the estimated mean yield of mixture m in environment l and $\hat{\mu}_{ml}^c = \sum_{i(m)} (\hat{\mu}_{il}/3)$ is the mean of the estimated mean yields of its three component varieties in pure stand in the given environment, as



b Variety Culma was not grown in these environments, hence mixing effect not available for Mix2

Table 2 A priori characteristics of each component variety (mixture membership indicated in parentheses) and the corresponding indices of inter-varietal diversity for each mixture (see text)

	Grain yield (1–5) ^b	Mildew (0–3)°	Leaf rust $(0-3)^{c}$	Net blotch $(0-3)^c$	Scald (0–3) ^c	Straw length (1–5) ^b	Weed suppressiveness (%)
Varietal characteri	stics						
Alabama (3)	3	0	1	2	2	1	19.2
Brazil (2, 4)	4	2	2	2	1	2	25.2
Cicero (2, 6)	3	0	2	1	2	2	23.9
Culma ^a (2)	_	_	_	_	_	_	_
Danuta (4)	3	0	1	2	2	5	49.0
Fabel (5, 6)	3	0	0	0	2	4	36.4
Harriot (5)	3	1	0	1	2	4	37.2
Landora (1)	3	0	2	1	0	4	29.5
Neruda (3)	3	0	2	3	2	3	27.7
Orthega (1, 4)	2	2	0	1	1	4	38.5
Otira (1)	4	0	2	2	2	2	31.3
Prestige (3)	3	0	1	3	1	3	27.6
Punto (6)	2	0	1	2	1	2	25.0
Sebastian (5)	4	3	1	2	1	2	18.9
Inter-varietal dive	rsity						
Mix1	1	1.16	1.16	0.58	1	1.16	4.8
Mix2 ^a	_	-	_	_	_	_	_
Mix3	0	0	0.58	0.58	0.58	1.16	4.9
Mix4	1	1.16	1	0.58	0.58	1.53	11.9
Mix5	0.58	1.53	0.58	1	0.58	1.16	10.3
Mix6	0.58	0	1	1	0.58	1.16	6.9

a No a priori information was available for the variety Culma

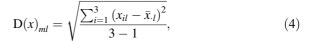
obtained from Eq. 1. Another measure of mixing effect, the extent to which mixtures were able to match the performance of the highest-yielding component variety in a given environment was found as

$$MEmax_{ml} = \frac{\hat{\mu}_l^m - \hat{\mu}_{ml}^{max}}{\hat{\mu}_{ml}^{max}},$$
 (3)

where $\hat{\mu}_{ml}^{\max}$ is the maximal estimated pure stand mean yield for any component variety of mixture m in environment l, as obtained from Eq. 1.

Inter-varietal diversity

In order to quantify for each mixture in each environment the inter-varietal diversity with respect to a number of plant characteristics, the standard deviation among component varieties for each characteristic was calculated as



where x_{il} is the value of component variety i in environment l and $\bar{x}_{\cdot l}$ is the mean value of all components of mixture m in that environment. A priori characteristics from external sources as well as pure stand observations in each of the 17 trials were obtained. Seven a priori characteristics on the potential performance of component varieties were derived from a national database (Table 2; for further details, see Østergård et al. 2008). These included index values for grain yield potential, susceptibility towards each of the four prevailing foliar diseases: powdery mildew ($Blumeria\ graminis$), leaf rust ($Puccinia\ hordei$), net blotch ($Pyrenophora\ teres$), and scald ($Rhynchosporium\ secalis$), and straw length potential measured under weed and disease free conditions. In addition, values



^b All growth potential indices: 1 (very low) to 5 (very high)

^c All disease susceptibility indices: 0 (resistant) to 3 (very susceptible)

on the potential weed suppressiveness (%) were obtained from Hansen et al. (2008), describing the average reduction in weed coverage in the plots of each variety compared to the 90% quantile based on all single-plot observations of that study.

A number of variety characteristics were observed in the studied environments, of which eight corresponded to the assessed a priori characteristics: harvested grain yield (at 85% dry matter), disease severity of each of the four prevailing foliar diseases (in % leaf area infected), infestation by tall annual weeds and creeping weeds, respectively (each in % ground cover; see Østergård et al. 2008), as well as straw length (after flowering, in cm). Assessments were detailed enough for predicting yield loss from a priori disease and weed competitiveness information, as shown by Østergård et al. (2008). Additionally, recorded dates of plant emergence, heading and maturity were used as variety characteristics (relative to the sowing date as number of days-until-emergence, number of days-until-heading, and number of daysuntil-maturity, respectively). Lodging was measured but the data were not appropriate for statistical analysis. All characteristics were observed for each genotype in each replicate, and genotypic means were estimated from models such as that in Eq. 1. Weed levels were not observed in the conventionally managed trials so these were all set to zero, assuming that the herbicide treatment was fully effective. The estimates of disease severity and ground cover of weeds for each genotype in each environment were third root transformed to optimize distributional properties prior to further analysis in analogy with previous analyses of the present data (cf Østergård et al. 2008).

Regression on inter-varietal diversity

For all a priori characteristics except grain yield potential, the effect of inter-varietal diversity on the yield of a given mixture in a given environment was likely to depend on biotic stresses, i.e. the environmental loads of the diseases or weeds of importance for the potential expression of the characteristic. Where relevant, estimates of inter-varietal diversity were therefore multiplied with the corresponding environmental load prior to regression in order to provide a weighted covariate. This procedure was similar to the methodology used by Østergård et al. (2008) for the prediction of variety characteristics. The loads applied

were 95% percentiles of all single plot observations in each environment (data from Østergård et al. 2008).

For most of the observed characteristics, actual environmental load was already part of the observation and the regression was done without environmental loads. For effects of inter-varietal diversity in observed straw length and phenological characteristics, however, a potential dependency on the environmental weed loads was hypothesized and correspondingly tested (see Table 3 for a list of all tested combinations).

The relationships between mixing effect and each type of inter-varietal diversity were analysed using a mixed model linear regression model of the general form

$$MErel_{ml} = \delta_{ml} + \beta_m * Z_{ml} + F_m + G_l + H_{ml}, \qquad (5)$$

where MErel_{ml} is obtained from Eq. 2, δ_{ml} is the expected mean mixing effect, β_m is the regression coefficient on Z_{ml} , and F_m , G_l and H_{ml} are random effects of mixture, environment and residual error, respectively. Depending on the trait, Z_{ml} equals the inter-varietal diversity $D(x)_{ml}$ or its product with environmental load L_l (as described above).

Estimation of parameters was done by log-likelihood maximization, and the effect of each type of intervarietal diversity was tested by likelihood ratio test against the intercept-only model ($\beta_m = 0$). Variation in mixing effect within each of the three management systems was as large as between them, e.g. the 'organic' environments ranked between third and sixteenth with respect to yield level. Therefore, genotypic performances were not compared between systems and instead the 17 environments were considered as representing large environmental variation.

The amount of variation explained by the covariate was assessed by a coefficient of determination, comparing total variance of the random effects in the covariate model with that in the intercept-only model as

$$R^{2} = 1 - \frac{\sum \hat{\sigma}_{(c)}^{2}}{\sum \hat{\sigma}_{(0)}^{2}}$$

$$= 1 - \frac{\sum \hat{\sigma}_{F(c)}^{2} + \sum \hat{\sigma}_{G(c)}^{2} + \sum \hat{\sigma}_{H(c)}^{2}}{\sum \hat{\sigma}_{F(0)}^{2} + \sum \hat{\sigma}_{G(0)}^{2} + \sum \hat{\sigma}_{H(0)}^{2}},$$
(6)

where (c) denotes variance components from the covariate model, (0) denotes variance components from the intercept-only (null) model, and letters F, G and H denote the variance components of the random terms in Eq. 5. Since total variance in the covariate model may become larger than that of



Table 3 Results of mixed model regression of mixing effect against each type of inter-varietal diversity, of which some characteristics are weighted by environmental loads (see text)

For each relationship are provided regression coefficient estimates (β , given in % point), test probabilities from likelihood ratio tests (P_{LRT}), and the coefficients of determination (R^2) a Negative regression coefficient estimates indicate that mixing effect decreases with increasing inter-varietal diversity b Probability levels are designated as bold (<5%) or

Inter-varietal diversity of	Environmental load	β (%) ^a	$P_{\mathrm{LRT}}^{}^{\mathrm{b}}}$	R^{2} (%)
A priori				
Grain yield potential	_	3.223	0.04	7.6
Mildew susceptibility	Mildew	0.009	0.80	0.1
Leaf rust susceptibility	Leaf rust	0.718	0.26	6.0
Net blotch susceptibility	Net blotch	0.031	0.78	0.3
Scald susceptibility	Scald	-0.012	0.96	0.0
Straw length potential	Tall weeds	0.022	0.39	1.4
Straw length potential	Creeping weeds	-0.010	0.53	0.7
Weed suppressiveness	Tall weeds	0.002	0.57	0.9
Weed suppressiveness	Creeping weeds	-0.002	0.36	0.7
Observed				
Grain yield	_	0.297	0.30	3.0
Mildew disease severity	_	0.058	0.96	0.0
Leaf rust disease severity	_	7.410	0.33	6.0
Net blotch disease severity	_	2.618	0.11	-0.4
Scald disease severity	_	4.257	0.16	1.7
Tall annual weeds infestation	_	-5.666	0.07	4.2
Creeping weeds infestation	_	1.712	0.51	1.0
Straw length	_	0.271	0.11	3.7
Straw length	Tall annual weeds	0.036	0.02	3.8
Straw length	Creeping weeds	0.012	0.16	-5.3
Days-to-heading	_	-0.049	0.89	0.1
Days-to-heading	Tall annual weeds	0.003	0.71	0.1
Days-to-heading	Creeping weeds	-0.003	0.65	0.4
Days-to-maturity	_	-0.256	0.55	0.4
Days-to-maturity	Tall annual weeds	0.006	0.67	0.1
Days-to-maturity	Creeping weeds	-0.008	0.20	1.9
Days-to-emergence	_	1.440	0.40	2.1
Days-to-emergence	Tall annual weeds	0.009	1.00	0.3
Days-to-emergence	Creeping weeds	0.008	0.79	-0.4

the intercept-only model, R^2 may become negative and is then interpreted as zero.

Yield stability across environments

Three concepts of genotype stability were applied. For static stability sensu Lin et al. (1986), an environmental variance was defined for each genotypic entity i as the variance of yields of genotypic entities across recorded trial environments:

$$s_i^2 = \frac{\sum_l (\hat{\mu}_{il} - \hat{\mu}_{i.})^2}{17 - 1},\tag{7}$$

where $\hat{\mu}_{il}$ is the estimated grain yield of genotypic entity *i* in trial environment *l* and $\hat{\mu}_{i}$ is the mean yield

of the genotypic entity across the 17 trial environments. For this measure, largest stability is seen at small s_i^2 values.

A dynamic measure of genotype stability (sensu Lin et al. 1986), describing the adaptability of genotypes (i.e. their responsiveness to environmental productivity), was obtained from the linear regression model

$$\hat{\mu}_{il} = a_i + b_i \cdot \hat{\mu}_{\cdot l} + d_{il}, \tag{8}$$

where a_i is the intercept, b_i is the regression coefficient of genotypic entity i, $\hat{\mu}_{.l}$ is the expected mean yield of all genotypic entities grown in trial environment l (Table 1), being used as the best available estimate of productivity in each environment, and d_{il} is used to denote error, being deviations from the fitted



italic (<10%)

regression line. Regression coefficients b_i then describe the ability of a genotypic entity to respond to environmental productivity, with an average coefficient of responsiveness corresponding to $b_i = 1$, following the average level of environmental interaction of all the varieties considered. A genotype with $b_i = 1$ is considered as possessing average responsiveness to environmental conditions (Finlay and Wilkinson 1963). Higher responsiveness is then found at $b_i > 1$, and lower responsiveness is found at $b_i < 1$. Hypotheses that $b_i = 1$ and $b_{i1} = b_{i2}$ were tested by t-tests.

The variance of deviations from the regression model in Eq. 8, $s(d_i)^2$, was used as a third measure of stability describing the sensitivity of genotypic entities to biotic and abiotic environmental factors apart from those defining general environmental productivity.

Results

Relative mixing effects

The average mixing effect was 1.9% (an average increase of 0.9 hkg ha⁻¹), which was significantly different from 0 (t-test, P < 0.001) for all mixtures across all trial environments, with lower and upper quartiles of mixing effect being -0.8 and 4.4%, respectively. Individual mixing effects ranged from -12.5 to 15.5% (Fig. 1a), varying significantly between mixtures (P < 0.05) and between trial environments (P < 0.05) when considering both as fixed effects in a simple analysis of variance. The average mixing effect was significantly positive in three of the six mixtures. Mix1 provided the highest average mixing effect of 4.0% (P < 0.001), corresponding to an average increase of 2.1 hkg ha⁻¹. The mixing effects of this mixture were almost exclusively positive (except a value of -0.3% in one environment), indicating that it yielded consistently more than the average of its components across all environments (Fig. 1a). Mix5 and Mix6 provided significant mixing effects of 2.4% (P < 0.05) and 2.9% (P < 0.01), corresponding to average increases of 1.3 and 1.2 hkg ha⁻¹, respectively (Table 4). The other mixtures provided insignificant average mixing effects. Given the incomplete use of management systems across locations and years, these factors were considered to be potentially confounded and could therefore not be tested.

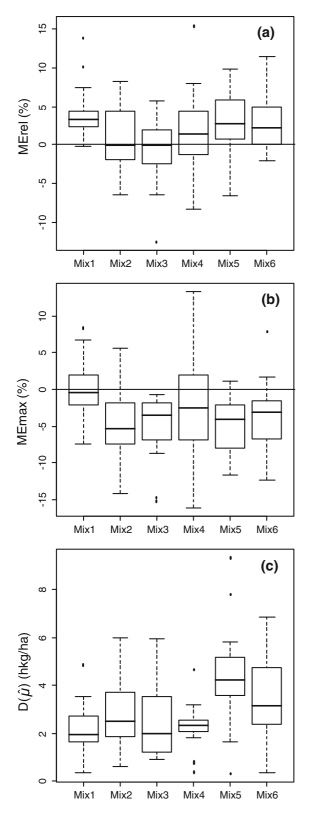
Effects of inter-varietal diversity

Only two of 28 tested linear relationships between inter-varietal diversity and mixing effect were significant (Table 3), namely (1) straw length weighted by load of tall annual weeds (P=0.02) and (2) grain yield potential (P=0.04). All but one of the disease-related characteristics (based on a priori as well as observed component variety characteristics) had positive but insignificant relationships to mixing effect, yet, inter-varietal diversity in leaf rust susceptibility and observed leaf rust severity both explained some variation in mixing effect ($R^2=6.0\%$ in both cases). No relationship with any inter-varietal diversity in phenology was found.

The first of the significant relationships showed that increased height diversity in the crop stand confers increased yields when competing with weeds of the tall annual type. A positive relationship with mixing effect was also suggested for inter-varietal diversity in straw length, although this relationship was not significant (Table 3). The relationship between mixing effect and the related (and oppositely directed) inter-varietal diversity in observed infestation by tall annual weeds was negative and almost significant (P = 0.07), suggesting that the yield advantage of growing varieties in mixtures was smaller when their ability to suppress these weeds (i.e. as observed in pure stand) was more diverse. The relationships between mixing effect and each of the corresponding types of inter-varietal diversity, straw length potential and weed suppressiveness, were far from significant (Table 3).

The second of the significant relationships showed that mixing effects were higher when mixing varieties more diverse in grain yield potential. Likewise, intervarietal diversity in observed grain yield, generally termed $D(\hat{\mu})$, had a positive relationship with mixing effect (Table 3). However, despite the overall increase in mixing effect with larger yield differences among component pure stand yields, as indicated also in Fig. 2a, the relationship was not significant (P =0.30). Mainly, Mix5 and Mix6 contributed to this putative relationship, whereas Mix 1 (with the largest mixing effect) did not (Fig. 2a). On the other hand, $D(\hat{\mu})$ was significantly related to mixture performance in terms of outyielding of component varieties, $MEmax_{ml}$. Overall, significantly higher values of MEmax_{ml} were found at lower values of $D(\hat{\mu})$ ($\beta_m =$ -0.013; P < 0.001; least squares regression). Hence,





◄ Fig. 1 Mixture-wise boxplots of **a** relative mixing effects, **b** mixture yield relative to highest yielding component, and **c** D($\hat{\mu}_i$). For each mixture is shown the median (*thick vertical lines*), lower and upper quartiles (*lower and upper ends of boxes*), standard deviations (*whiskers*), and extreme values (*black dots*) among the 17 environments

mixtures were generally more likely to outyield the highest yielding component (MEmax $_{ml}$ > 0) when the components had more similar yields (low values of $D(\hat{\mu})$; see Fig. 2b). Mix1 provided a higher yield than (outyielded) all of its component varieties in 8 environments, whereas for the other mixtures this effect was much less pronounced, occurring in only 3, 0, 5, 1 and 3 of the environments, respectively (Figs. 1b, 2b). As seen from the lower values of $D(\hat{\mu})$ (Fig. 1c), the component varieties in each of mixtures 1, 3 and 4 generally had more similar yields in each environment than the component varieties of Mix5 and Mix6 (all P < 0.05 in pairwise t-test with adjustment for multiple comparisons) whereas values of Mix2 were not different from those in any of the other mixtures (Fig. 1c). The $D(\hat{\mu})$ of Mix4 were particularly similar across growth environments, as seen from the small inter-quartile range.

Yield and stability analysis

Average grain yield levels ranged from 48.6 to 53.7 hkg ha^{-1} for mixtures and from 45.1 to52.1 hkg ha⁻¹ for component varieties in pure stand (Table 4). Among all genotypic entities, the highest average yield level was found in Mix1 and the lowest in the variety Fabel. It is seen by plotting grain yields of each mixture and its component varieties separately against average environment yields (Table 1; Fig. 3) that Mix1 outyielded its components in the four environments with the highest levels of productivity (Fig. 3a). One of the component varieties (Harriot) of Mix5 yielded generally higher than the mixture and the other components across the range of environmental productivity (open squares in Fig. 3e); yet, under higher levels of environmental productivity the yield of Mix5 approached that of Harriot and occasionally exceeded it. Mix6 yielded generally well, having higher yields than each of its components in pure stand under less productive conditions (Fig. 3f). In only one mixtureenvironment combination did the mixture provide a considerably lower yield than all of its components (Mix3, with a yield of app. 35 hkg ha⁻¹; Fig. 3c).



Table 4 Grain yield level $(\hat{\mu}_i)$ and measures of yield stability $(s_i^2, b_i, s(d_i)^2)$ of each component variety and variety mixture across environments

	$\hat{\mu}_i$ (hkg ha ⁻¹)	$s_i^2 \text{ (hkg}^2 \text{ ha}^{-2}\text{)}$	b_i	$s(d_i)^2 (hkg^2 ha^{-2})$
Varieties ^a				
Alabama (3)	48.0	5.67	1.07	3.45
Brazil (2,4)	49.9	5.00	1.03	2.55
Cicero (2,6)	50.0	6.24	1.15	3.24
Culma (2)	47.9	5.61	0.98	2.16
Danuta (4)	52.0	5.42	1.08	2.25
Fabel (5,6)	45.1	3.68	0.85	3.58
Harriot (5)	52.1	4.88	1.02	2.53
Landora (1)	51.9	6.79	1.18	3.07
Neruda (3)	50.6	5.35	1.06	2.90
Orthega (1,4)	51.3	5.25	1.06	2.51
Otira (1)	51.6	4.63	0.99	2.27
Prestige (3)	49.2	5.05	1.05	1.71
Punto (6)	47.1	4.66	1.01	1.52
Sebastian (5)	48.6	5.06	1.00	3.14
Varieties mean	49.9	5.21	1.04	2.60
Mixtures ^b				
Mix1	53.7	5.90	1.13	1.87
	(51.6)	(5.56)	(1.08)	(2.62)
Mix2	49.8	6.19	1.10	1.61
	(49.3)	(5.62)	(1.05)	(2.65)
Mix3	49.0	5.64	1.11	2.04
	(49.3)	(5.36)	(1.06)	(2.69)
Mix4	52.0	6.31	1.13	3.22
	(51.1)	(5.22)	(1.06)	(2.44)
Mix5	49.8	5.04	1.04	2.10
	(48.6)	(4.54)	(0.96)	(3.08)
Mix6	48.6	4.05	0.94	1.69
	(47.4)	(4.86)	(1.00)	(2.78)
Mixtures mean	50.4	5.49	1.08	2.06

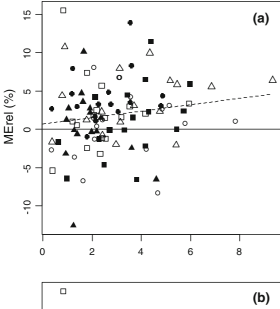
^a Mixture membership of varieties provided in parentheses

In the following, the three stability estimates of each genotypic entity are considered (Table 4). First, the environmental variance, s_i^2 , of all mixtures except one (Mix4) was within the range of their component varieties. The environmental variance of the variety Fabel was the lowest of any of the considered genotypes. Second, for most mixtures and component varieties the responsiveness to environmental productivity was higher than the average of all genotypes grown in the field trials (as indicated by regression coefficients b_i larger than 1). The regression coefficients of five mixtures were non-significantly larger than the average coefficients of their components and non-significantly larger than 1 (t-tests; not shown). The regression coefficient of Mix6 was smaller than

the average of its components by 0.06 and numerically smaller than 1, only being higher than its component Fabel. Fabel furthermore had the lowest average yield among genotypic entities, indicating that this variety had a relatively higher adaptability to low-yielding environments (as seen from the lower left-most position in Fig. 4). Conversely, Mix1 and its component Landora showed the highest levels of responsiveness to high environmental productivity (as seen from their upper-right-most position in Fig. 4). Third, the variance of deviations from the regression line, $s(d_i)^2$, was generally lower for mixtures than for pure stand varieties, indicating that mixtures were less sensitive to biotic and abiotic stresses. One of the mixtures (Mix4) was significantly more sensitive than



^b Component means in italics and parentheses below each mixture value



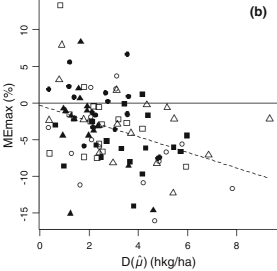
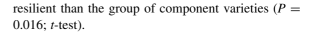


Fig. 2 Linear regressions of **a** relative mixing effect and **b** mixture yield relative to highest yielding component against the standard deviation of component variety yields in each environment. *Individual data points* are shown for Mix 1 (*full circles*), Mix 2 (*full squares*), Mix 3 (*full triangles*), Mix 4 (*open circles*), Mix 5 (*open squares*), and Mix 6 (*open triangles*)

any of the remaining mixtures and most of the pure stand varieties (P < 0.001; z-test), as seen in Fig. 3d. The variety Punto (component of Mix6) showed the lowest level of sensitivity of all genotypic entities, including mixtures, whereas the variety Fabel (component of Mix5 and Mix6) showed the very highest level of sensitivity. When disregarding the latter, the group of mixtures was found to be significantly more



Discussion

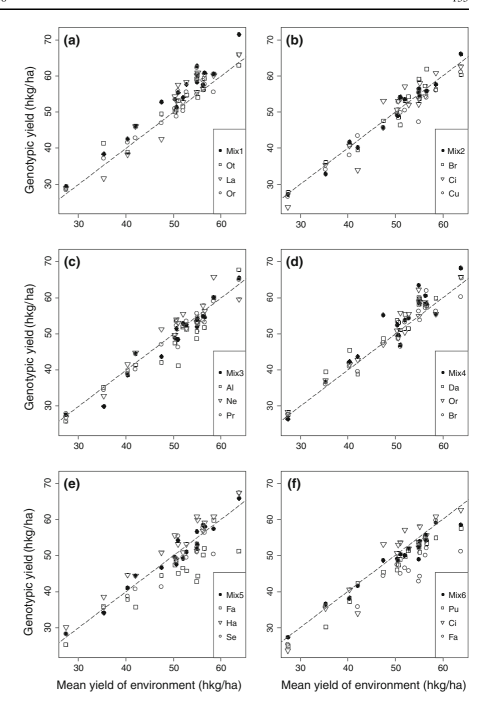
Effects of inter-varietal diversity

Among all types of inter-varietal diversity, yield potential diversity had the largest influence on mixing effect (i.e. largest coefficient of determination, Table 3), thereby confirming previous findings from a literature survey on variety mixtures of wheat and barley (Kiær et al. 2009). Furthermore, the relative mixing effect was generally higher among varieties of more diverse height and more so in the presence of tall annual weeds. This supports the hypothesis that mixtures of varieties of varying straw length have an actual advantage in terms of weed suppression (Kaut et al. 2009). Interestingly, mixing effects tended to be smaller when the level of suppression of tall annual weeds by component varieties was more diverse. This indicated that the yield advantage of mixing varieties is lower if one component is particularly effective or particular poor at suppressing these weeds.

Few types of inter-varietal diversity were thus correlated with mixing effect, even when accounting for relevant environmental loads (Table 3). The finding that most mixtures (all except Mix4) were less sensitive to biotic and abiotic environmental stresses than most of their component varieties suggests the contrary. A number of possible explanations are therefore worthwhile considering. First, grain yield is the ultimate result of multiple genotype-environment interactions, and differences in harvested grain yield can therefore be seen as a composite descriptor of the complex inter-varietal diversity that is insufficiently described by single factors. Hence, mixing effect could be the result of many small effects, each of which was too small to detect with the current analysis and experimental set-up. Second, the observed variation in mixing effect could be strongly influenced by characteristics and factors other than those considered. As an example, below-ground characteristics and interactions are often of greater importance for plant performance than those above ground (e.g. Wilson 1988) but are usually difficult to observe; root length and root biomass of the component varieties grown hydroponically were found to have a positive relationship with



Fig. 3 Grain yields of mixtures (black dots) and their component varieties (open triangles, open circles and open squares) plotted against environmental mean yields (a Mix1; b Mix2 etc.). Component names provided as the first two letters in each name. Broken lines indicate average stability (i.e. regression line with a = 0, b = 1)



mixing effect (pers. comm., N.-O. Bertholdsson). Third, whereas most of the trials in the present study were organic or low-input, the a priori characteristics used were based on VCU tests under conventional cultivation and may not be optimal for describing interactions in the studied environments. Last, some of

the observed varietal characteristics may have been biased by interplot interference, as discussed below.

Inter-varietal diversity in disease resistance genes is well documented as an effective means of controlling fungal pathogens and stabilizing yield under disease in variety mixtures (Smithson and Lenné 1996; Finckh



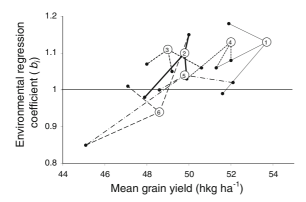
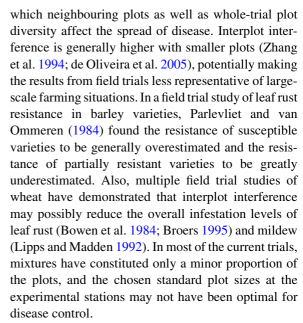


Fig. 4 Average grain yields plotted against environmental regression coefficients of variety mixtures 1–6 (numbered circles) and their component varieties (black dots), respectively (connected by lines)

et al. 2000; Newton et al. 2002). In practice, the largest proportion of commercial variety mixtures is grown for the purpose of fungal disease control. In Denmark, official certification procedures for variety mixtures set minimum requirements with respect to the disease resistance of component varieties (Østergård and Jensen 2005). The mixtures were selected to follow these requirements, which implied relatively little inter-varietal diversity in disease susceptibility and severity (not shown), which in turn may have been the cause of the low correlation with mixing effect. A number of other circumstances may have contributed to this lack of significant relationships. First, the applied disease susceptibility scores (0-3) may be too simple to be used as predictors of disease development in mixtures, e.g. two component varieties may have the same low susceptibility score due to different resistance genes, so that in a specific environment only one of the components may be resistant to the actual pathogen strain; the weighting by environmental disease load would not be able to compensate for this. Second, simultaneous infection by multiple pathogens may have had a non-additive yield effect. As an example, when all traits in the full data set were combined and weighted with observed disease loads of powdery mildew, leaf rust and net blotch, a nonadditive effect of powdery mildew and leaf rust on grain yield of varieties was found (Østergård et al. 2008). In the present study, the actual data were considered too sparse for such an approach. Last, the observed disease levels may have been biased by interplot interference, a well-studied phenomenon in



Interplot interference from competition between neighbouring plots are found to introduce a similar type of error. Hence, in field trials of cereal varieties of different height, interplot interference can result in shorter varieties being depressed due to shading from taller neighbours, while the yields of taller varieties are increased (e.g. Kempton 1982, Kempton and Lockwood 1984). Interplot interference is found to be higher in trial environments of higher fertility (e.g. Aastveit et al. 1989; Clarke et al. 1998). Varying levels of interference across environments may thereby decrease the ability to detect any overall relationship in multi-environment trials such as that presented. The performance of nearest neighbour plots has previously been modelled in order to mitigate or eliminate the effects of interference (e.g. Kempton and Howes 1981; Kempton 1985; Talbot et al. 1995; Durban et al. 2001). However, this may sometimes introduce larger bias than that of the interference itself (Ainsley et al. 1995) and such models were therefore not applied.

Yield and stability analysis

Considering mixing effect on grain yield as the joint result of many types of inter-varietal and genotypeenvironments interaction, the results discussed above suggest that each type had only a minor influence. In trials such as these, where mixtures were not designed to study specific types of inter-varietal interaction, it may be more fruitful to consider general genotypic



responses such as adaptability and resilience across the range of environmental conditions. These stability concepts are frequently used to compare crop properties (for a review, see Piepho 1998), including comparisons of variety mixtures and their components (e.g. Smithson and Lenné 1996; Juskiw et al. 2001; Cowger and Weisz 2008). Here, the overall yield variation, of mixtures was slightly higher than that of pure stand varieties, and since this measure encompasses both the response to environmental productivity and sensitivity to weeds and diseases, it seems more informative to consider these sources of variation separately.

Comparisons of genotypic responsiveness to environmental productivity are commonly obtained through the use of regression coefficients, b_i (i.e. adaptability; Yates and Cochran 1938; Finlay and Wilkinson 1963; Eberhart and Russell 1966; Nurminiemi et al. 1996). In order to promote genotype comparisons across different studies, values of b_i should always be interpreted in conjunction with the overall yield level of the genotype (Piepho 1998). In essence, genotypes with $b_i > 1$ are responsive to high environmental productivity when having a high yield level (Clay and Allard 1969; Juskiw et al. 2001), such as Landora and Mix1. Oppositely, in conjunction with a low yield level it is indicative of high vulnerability to low environmental productivity, of which none were found in this study (Fig. 4). We found that b_i values were sometimes higher than that of all their components (Mix3 and Mix4), whereas Juskiw et al. (2001) by a comparable approach found b_i values of mixtures to be intermediate relative to component values. In the review of 36 studies, Smithson and Lenné (1996) found the environmental regression coefficients of both mixtures and component varieties to be very variable between studies. It is important to keep in mind that b_i depends directly on the considered set of genotypes and environments. Absolute values of b_i are therefore interpretable mainly within the considered data set, as has been shown for Nordic barley varieties (Nurminiemi and Rognli 1996), but ratios of b_i of mixtures to component averages may be used for comparison between similar studies.

Variety mixtures are generally seen to deviate less from the environmental regression line than their component varieties (Smithson and Lenné 1996). This was confirmed here, using $s(d_i)^2$ as an overall indicator of sensitivity. All mixtures except Mix4 were more resilient than two or all of their component varieties

(Table 4), Mix4 being among the least resilient of all mixtures and component varieties. Compared to the other mixtures, the component varieties of Mix4 were expected (from a priori characteristics) to differ the most in straw length and weed suppressiveness (Table 2), yet it remains unsolved as to how this may have contributed the observed sensitivity of Mix4.

Interaction between component varieties

The identified differences in stability of mixtures can be attributed to different levels of complementarity and compensation among component varieties. Since yields of single component varieties within mixture plots were unavailable, this interpretation can only be tentative.

Component variation in the environmental regression coefficient, b_i , may be an important indicator of the ability of component varieties to compensate for lower yields of other varieties in the mixture. Since compensation can occur at high as well as low levels of environmental productivity, varieties of lower responsiveness will be able to compensate for more responsive genotypes in less-productive environments and vice versa. It seems that the potential for compensation among mixture components is larger the more the components differ in response to environmental productivity measured by b_i . A simple regression of six data points (one for each mixture) supported this hypothesis, showing that the average mixing effect across environments increased significantly with the standard deviation of b_i among component varieties (r = 0.64). In Mix1, the component variety Orthega, (b_i close to the average of component varieties), was seen to have compensated the variety Landora (highest b_i of any genotypic entity) at lower environmental productivity, whereas the opposite was seen at higher environmental productivity (Fig. 3a). The lowest b_i of any genotypic entity was seen in the variety Fabel, which was bred for cultivation under low-input organic conditions. Fabel was component of Mix6, the b_i of which was also lower than 1 and markedly lower than those of the remaining mixtures. This was not because the other component varieties could not compensate fully for the generally low yield of Fabel, but rather, it was a result of higher compensation in less productive environments (Fig. 3f). In the perspective that varieties can thus differ (and compensation occur) at all levels of productivity, the previously



stated hypothesis that mixing effect will be higher when a compensating component variety has a steeper response to environmental productivity (e.g. Stützel and Aufhammer 1990) may seem over-simplistic.

Compensation will tend to be lower in mixtures whose component varieties provide more similar grain yields, i.e. when $D(\hat{\mu})$ is smaller. This was observed in Mix5 and Mix6, where Fabel provided lower yields in most trials compared to the other components. Additionally, one of the components of Mix5 (Harriot) generally yielded more than the other components throughout the range of productivity. The overall relative mixing effect of this mixture was thus probably due to compensation by Harriot through uptake of the resources not utilized by Fabel and Sebastian. Positive effects of inter-varietal complementarity, will thus tend to be more important for overall mixing effect with smaller $D(\hat{\mu})$. In this case, mixture yields above the level of the highest yielding component (MEmax $_{ml} > 0$) would clearly indicate an effect of complementarity. Substantially more examples of outyielding were seen at lower $D(\hat{\mu})$ (Fig. 2b) and a significantly negative relationship between MEmax_{ml} and D($\hat{\mu}$) was confirmed, indicating that complementarity among mixture components is more important for mixing effect the less they differ in grain yield. Another support of this hypothesis was found in the mixture with the highest average mixing effect (Mix1). The component varieties of this mixture provided the lowest average $D(\hat{\mu})$ (Fig. 1c) and the highest average mixing effect (Fig. 1a) of all mixtures and actually produced higher grain yields than its best yielding component in nearly half of the trials, most likely deriving from inter-varietal complementarity.

Perspectives

The indication that the potential for compensation among mixture components is higher the more they differ in yield level may seem trivial but nonetheless highlights the divergence of focus (on variety mixtures) between researchers and farmers. A large mixing effect (in relative as well as absolute terms) may easily be obtained by including one or more mixture components that perform substantially worse than the remaining components. This may provide the opportunity to study compensation and other mechanisms between genotypes in a mixture; however, it may not be of agronomical interest as it would most likely be

accompanied by grain yield levels lower than what could be obtained using the best yielding variety. As an example, the requirement by farmers and authorities for a low level of disease susceptibility in all component varieties of marketed variety mixtures is contrasted by the fact that most published studies of fungal disease in variety mixtures involve one or more susceptible varieties (for reviews, see Smithson and Lenné 1996; Finckh et al. 2000).

The general experimental evidence of more stable yields of mixtures than the average of their pure stand component varieties may justify large-scale cultivation of variety mixtures in variable environments, such as organic farm systems. However, mixtures would be even more advantageous to farmers in general if, in addition, mixture yields were comparable to the highest yields of component varieties. Such mixtures exist, as exemplified by Mix1 which was exceptionally good with respect to both yield level and stability. The component varieties of Mix1 were all relatively highyielding. Noteworthy, a dynamic reference mixture composed of the three most high-yielding varieties from the previous year has been used at reference standard in Danish VCU testing for a number of years as it provides not only more stable but often also higher yields than any pure stand variety (pers. comm., M. Haastrup).

In essence, the successful formation of variety mixtures depends on the ability to select optimal a priori combinations of variety characteristics. The reported study suggests that, for a number of possible reasons, mixing effect may be difficult to predict from inter-varietal diversity with respect to single quantitative characteristics, even in the rare cases when a priori information on environmental conditions such as disease and weed loads is available. Proposed methods for predicting the best variety combinations based on thorough field trials, e.g. considering disease resistance (Mundt et al. 1995; Newton et al. 2008) or general mixing ability (Valentine 1982; Gallandt et al. 2001), may not always be feasible for practical farming. Prominently, new varieties often enter and leave the market within a short time-scale so that seeds of the varieties tested in such trials may be unavailable when recommendations based on those trials are eventually finished. Our findings suggest that deliberate combination of more general variety characteristics such as high yield levels and different environmental responsiveness is more attractive from an agronomic



perspective, conferring a high potential for interaction between component varieties and their environment. If variety mixtures are to be adopted more widely by farmers, future research should address such mixtures more deliberately.

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