A comparison between Joint Regression Analysis and the Additive Main and Multiplicative Interaction model: the robustness with increasing amounts of missing data

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ABSTRACT: This paper joins the main properties of joint regression analysis (JRA), a model based on the Finlay-Wilkinson regression to analyse multi-environment trials, and of the additive main effects and multiplicative interaction (AMMI) model. The study compares JRA and AMMI with particular focus on robustness with increasing amounts of randomly selected missing data. The application is made using a data set from a breeding program of durum wheat (Triticum turgidum L., Durum Group) conducted in Portugal. The results of the two models result in similar dominant cultivars (JRA) and winner of mega-environments (AMMI) for the same environments. However, JRA had more stable results with the increase in the incidence rates of missing values.

Keywords: AMMI models, genotype by environment interaction, joint regression analysis, missing values, durum wheat

Introduction

Joint Regression Analysis (JRA) has been widely used in crop sciences, to structure and understand Genotype by Environment Interaction (GEI) (Eberhart and Russell, 1966; Finlay and Wilkinson, 1963; Gusmão, 1985; Mooers, 1921; Pereira and Mexia, 2008; Yates and Cochran, 1938; Zheng et al., 2009), and in genetics, to analyse quantitative trait loci (QTL) by environment interaction (Emebiri and Moody, 2006; Korol et al., 1998). In this paper we are mainly interested in the approach proposed by Gusmão (1985) in which the precision in analysing series of randomized block experiments was highly increased, by considering environmental indexes for individual blocks instead of only one environmental index per experiment. In the literature some variants of JRA are also denoted as SREG (Sites Regression) model (Cornelius et al., 1992; Crossa et al., 2002; Setimela et al., 2007).

Williams (1952), Gollob (1968), Mandel (1971), Bradu and Gabriel (1978) and Gauch (1988) have made an important contribution to the development of additive main effects and multiplicative interaction (AMMI) models. These models have been widely used to analyze multi-environment trials because of their flexibility in allowing the use of several multiplicative terms to explain the GEI. One of the difficulties in choosing the right tool to analyse multi-environment trials arises when there are missing values in the two-way table of genotypes and environments. These missing values can be either systematic (Calinski et al., 1992; Denis and Baril, 1992), or selected completely at random in the two-way table.

This paper brings together the main features of JRA and AMMI models, and compares them for analyzing a durum wheat (Triticum turgidum L., Durum Group) trial with particular focus on robustness with increasing amounts of random missing data, either missing replications or missing cells (more likely when the proportion of missing values is high). The aim here is not to compare the method’s ability to estimate missing values in comparison to real data (Alarcón et al., 2010; Bergamo et al., 2008) but to compare the overall stability when increasing the incidence rate of missing values. An emphasis is made in the comparison between (i) the upper contour of JRA and the mega-environments of the AMMI model; and (ii) the stability of the dominant/winner genotypes across environments. To obtain the results for the JRA we developed an R code, and the MATMODEL software (Gauch and Furnas, 1991) was used to fit the AMMI models.

Materials and Methods

Joint regression analysis

JRA has proven to be an important model for analysing and interpreting the GEI of two-way classified data tables and continues to be largely used as a complement of traditional statistical analysis in genetics, plant breeding, and agronomy, for determining yield stability of different genotypes or agronomic treatments across environments (Crossa, 1990). JRA may also be used for the analysis of series of experiments in genotype comparison and selection. This technique is based on the adjustment of a linear regression, per genotype, of the yield on a synthetic variable measuring productivity, the environmental index.

JRA, when applied to two-way tables obtained from multi-environment trials, aims to determine the stability of the genotypes or agronomic treatments over a wide range of environmental conditions and to interpret the interaction (non-additivity). Let $Y_{ij}$ be a continuous response variable (usually yield) corresponding to a row factor $j, j = 1, ..., J$ (usually the genotypes), and a column factor $j, j = 1, ..., J$ (usually the...
environments. The model used for the analysis of multi-environment variety trials can be defined as

\[ Y_{ij} = \mu + G_i + E_j + GE_{ij} + \varepsilon_{ij} \]  

(1)

where \( \mu \) is the grand mean, \( G_i \) and \( E_j \) are the genotype and environment main effects, \( GE_{ij} \) is the interaction and \( \varepsilon_{ij} \) is the residual. A sub-model of (1), aiming at estimating some stability parameters for making comparisons between varieties is given by JRA, and allows us to partitioning the GEI into two parts of interest, i.e.

\[ (GE_{ij}) = b_i E_j + \delta_{ij} \]  

(2)

where \( b_i \) is a linear regression coefficient for the \( i \)-th genotype and \( \delta_{ij} \) a deviation (unexplained GEI) (Freeman, 1973). The JRA model can then be written as

\[ Y_{ij} = [\mu + G_i] + [E_j + b_i E_j] + \varepsilon_{ij} = [\mu + G_i] + [b_i E_j] + \varepsilon'_{ij}, \]  

(3)

where \( \varepsilon'_{ij} \) comprises both the unexplained GEI and the experimental error (Shukla, 1972). We assume fixed genotypic and environmental effects and random residual term.

The model (3) used in the present paper does not take into account the block effects since it uses the blocks as environments, following Gusmão (1985). If an experiment is designed with randomized blocks and the treatments correspond to the \( J \) genotypes to be compared, for each block in each design, the environmental index is measured by the average yield. For each of the \( J \) genotypes, a linear regression of yield on environmental indexes is adjusted.

**L2 environmental indexes**

For convenience, let us consider the joint regression model of the second equation in (3), where \( G_i' = \mu + G_i \), \( b_i' = 1 + b_i \), \( E_j \) is the environmental index corresponding to blocks instead of environments, \( b \) the number of blocks, \( Y_{ij} \) is a continuous response (e.g. yield) for cultivar/genotype \( i \) in block \( j \) if present, and the pairs \((G_i', b_i') \), \( i = 1, ... , I \), are the regression coefficients, for the \( I \) genotypes.

To obtain the estimates for the regression coefficients and the environmental indexes, the goal function to be minimized should be

\[ S(G', b', E) = \sum_{i=1}^{I} \sum_{j=1}^{b} p_{ij} (Y_{ij} - G_i' - b_i' E_j)^2. \]  

(4)

Usually the weight \( p_{ij} \) is 1 [0] when genotype \( i \) is present [absent] in block \( j \). These weights may differ from block to block to express differences in representativeness of the blocks and thus we take \( p_{ij} = p_j \) when the \( i \)-th genotype is present. The main problem in such modeling is how to estimate the parameters. However, the lately proposed so called zigzag algorithm (Pereira and Mexia, 2010) is very efficient in finding the estimates of \((G_i', b_i') \), \( i = 1, ... , I \), and \( E_j \). This zigzag algorithm is an alternating least squares based algorithm (Calinski et al., 1992; Denis and Baril, 1992; Digby, 1979; Gabriel and Zamir, 1979; Gauch and Zobel, 1990). For the complete case, Pereira and Mexia (2010) presented an alternative algorithm, the double minimization algorithm, which converges to the absolute minimum of the goal function (4) and is an adaptation of the algorithm first presented by Fisher and Mackenzie (1923). More details on the zigzag and double minimization algorithms can be found in Pereira and Mexia (2010).

**Upper contour**

When two of the regressions on genotypes intersect it means that one of the genotypes is better for higher environmental indexes while the other is preferable for lower environmental indexes. The intersection of regressions shows more than one genotype with similar performance. The upper contour
of the JRA is a concave polygonal (Mexia et al., 1997), constituted by segments of the adjusted regression lines, that contains the higher adjusted yields for the environmental indexes (Figure 1). Each of these segments will correspond to a range of variation of the environmental indexes in which the associated genotype will have the maximum adjusted yield (Pereira and Mexia, 2008). These genotypes are called dominant and should be selected. The remaining genotypes should be compared with the dominant to check whether they are dominated on entire range for the adjusted environmental indexes, [c, d]. If so, they can be safely discarded from the breeding program.

An analogy can be made between Figure 1 in this paper and Figure 2 in Gauch (1997), where the AMMI1 nominal yields for a corn trial is depicted as a function of the environment interaction principal component (IPC) axis 1. A more detailed comparison in what concerns the winner genotypes across the environments is presented latter in this paper.

Genotype comparison and selection

Let L be the number of dominant genotypes with dominant ranges \( c_i = \bar{\beta}_i, d_i = \bar{\beta}_i \), \( i = 1, \ldots, L \). The entire range for the environmental indexes will be \( c = \bar{\beta}_i, d = \bar{\beta}_i \). To have interaction between genotypes and environments there are two possible cases for different slopes, \( \bar{\beta}_i < \bar{\beta}_j \) and \( \bar{\beta}_i > \bar{\beta}_j \). After establishing the upper (dominant) non-dominant genotypes should be compared with the dominant ones. This comparison should be made on the left (right) extreme of the dominance range if the non-dominated genotypes have lower (greater) slope than the dominant one. So, when \( \bar{\beta}_i < \bar{\beta}_j \) \( \bar{\beta}_i > \bar{\beta}_j \) we are led to compare the adjusted values \( \bar{a}_i + \bar{\beta}_j \bar{x} \) and \( \bar{a}_j + \bar{\beta}_i \bar{x} \) at the environmental index \( \bar{\theta} \). These comparisons between slopes may be made using one of the following statistical tools: (i) one-sided \( t \) tests for the null hypothesis; (ii) Scheffé multiple comparison tests (Scheffé, 1959); (iii) Bonferroni multiple comparison method (Seber and Lee, 2003); (iv) Tukey multiple comparison method (only for the complete case); and (v) Control of False Discovery Rate which is robust against erroneous rejections (Benjamini and Hochberg, 1995). More details of these tests can be found in Pereira and Mexia (2008).

AMMI models

The core idea of the AMMI models is: (i) first apply the additive analysis of the variance model (ANOVA) to a two-way data (in the present case with genotypes and environments); and (ii) secondly apply the multiplicative principal component analysis (PCA) model to the residual from the additive model (in this case to the interaction) (Gauch, 1992). The AMMI model with \( N \) multiplicative terms can be written as

\[
Y_{ij} = \mu + \alpha_i + \beta_j + \sum_{n=1}^{N} \lambda_n \gamma_{ni} \delta_{nj} + \theta_{ir},
\]

where \( Y_{ij} \) is the yield of genotype \( i \) in environment \( j \); \( \mu \) the grand mean; \( \alpha \) the genotype main effects (deviations the genotype means minus the grand mean); \( \beta \) the environmental main deviations; \( \lambda_n \) the singular value for the PCA axis \( n \); \( \gamma_{ni} \) and \( \delta_{nj} \) are the genotype and environment and PCA scores for PCA axis \( n \); \( N \) is the number of PCA axes retained by the model; \( \theta \) the residual. If the experiment is replicated, an error term \( \varepsilon_{ijr} \), which is the difference between the \( Y_{ij} \) mean and the single observation for replicate \( r \), should be added.

The main purposes of the AMMI models were pointed out by Crossa (1990): (i) model diagnosis (Bradu and Gabriel, 1978); (ii) to clarify GEI (Crossa et al., 1990; Zobel et al., 1988); and (iii) to improve the accuracy of yield estimates (Crossa et al., 1990; Zobel et al., 1988).

Durum Wheat Yield Data

All the properties and comparisons presented in this paper are illustrated with a data set resulting from a breeding program in Portugal, carried out by the Portuguese National Plant Breeding Station (ENMP, Elvas) in the years of 1992/1993 and 1993/1994. It contains the yield from nine genotypes (CELTA; HELVIO; TE9006; TE9007; TE9008; TE9110; TE9115; TE9204; and TROVADOR) of durum wheat (Triticum turgidum L., Durum Group), measured in 11 environments (Benavila1; Revilheira; Evora; Elvas1; Beja1; Tavira1; Elvas2; Tavira2; Elvas3; Benavila2 and Beja2), and performed in complete random blocks with four replicates. These environments were obtained in two years, the first 6 in the first and the second 5 in the second year. Only the locations Tavira, Benavila and Beja were the same in both years. All the locations in this data set are in south Portugal, Tavira being at the sea side (Algarve) while the remaining in the inland (Alentejo). More details about this data set can be found in Pereira and Mexia (2010).

Simulation of missing values

Since the plants may be destroyed by animals, floods or during the harvest, and the yield measurements may be erroneously performed and inadequately introduced in the data base, missing values are common in agricultural experiments. When dealing with missing values researchers should decide between: (i) find a good tool to estimate the missing values (Alarcón et al., 2010; Bergamo et al., 2008), or (ii) chose a robust technique against missing observations to perform the analysis. In the present study we will be interested in the second approach, namely to compare the robustness of JRA and AMMI with the increasing of missing data. Our interest here is to study the case where the missing values were selected “completely” at random, instead of having systematic patterns (Calinski et al., 1992; Denis and Baril, 1992).

Our simulation procedure can be summarized in the following steps:

(i) Choose the incidence rate of missing values \( \alpha \) (e.g. \( \alpha = 5, 10, 25, 50, 75 \% \));

(ii) Remove, randomly, \( \alpha \% \) of the two-way table with genotypes and environments, leaving at least one observation in each environment and in each genotype;

(iii) to improve the accuracy of yield estimates (Crossa et al., 1990; Zobel et al., 1988).
In this case an Expectation-Maximization (EM) algorithm provides an effective general strategy for obtaining maximum likelihood estimates (Gauch, 1992). This procedure has been adapted for AMMI and is called EM-AMMI (Gauch and Zobel, 1990), and is implemented in the MATMODEL software (Gauch and Furnas, 1991).

Results and Discussion

A comparison between the algorithms and the alternative methods

This subsection presents a comparison between the two algorithms mentioned in the above section - (i) zigzag algorithm (Pereira and Mexia, 2010) and (ii) double minimization algorithm (Pereira and Mexia, 2010); and the two methods based in the joint regression model - (iii) the regression analysis of the mean yield of individual genotypes on the overall mean of the trial (Finlay and Wilkinson, 1963), and (iv) the regression analysis of the genotype mean yield on block mean, proposed by Gusmão (1985). This comparison is illustrated with a numerical example using the durum wheat yield population. Estimates of intercept, slope and the coefficients of determination, obtained from the Finlay and Wilkinson (1963) and Gusmão (1985) methods, and the zigzag and double minimization algorithms are presented in Table 1.

To compare these four procedures it is important to analyze the slopes and coefficients of determination. They produced almost the same results regarding the ordering of the genotypes per slope (only the Gusmão’s method gave a small difference). The coefficients of determination are mainly similar, the zigzag and Double Minimization algorithms being lower than Gusmão (1985) only for three environments (HELVIO, TE9110 and TE9115). Moreover, the zigzag and double minimization have completely agreed and may be seen as the most suited for regression analysis of complete randomized blocks because of their convergence to the minimum of the loss function (4).

Another comparison can be made regarding the sums of the sums of squares of residuals for the two procedures and two algorithms (Table 2). Here the advantage of the zigzag and double minimization algorithms over the two other procedures is evident since the algorithms induce lower sums of the sums of squares of residues. This result is true for all the examples and the mathematical proof can be found in Pereira and Mexia (2010). If we compute the pairwise Pearson correlations between the environmental indexes for the four alternatives in Table 2, we conclude that all the obtained environmental indexes are highly correlated (minimum of 0.984). In particular, the results obtained using the zigzag and double minimization algorithms have a coefficient of correlation of 1.000 since they completely agree with each other, and they are slightly better than the Finlay and Wilkinson (1963) and Gusmão (1985) approaches. In the case of a comparison using α-designs or incomplete blocks (instead of the randomized complete block design) some advantage within the two algorithms could better be presented (Pereira and Mexia, 2010).

Genotype comparison and selection

The results for some of the multiple comparison tests mentioned above can be found in Table 3. The graphical representation of the dominant genotypes, together with the ranges of dominance (i.e. the lower and upper bound for the interval where the each genotype is dominant) and environments where that dominance occurs, is depicted in Figure 1. The bounds of the environmental indexes 2.21 and 8.84 (Table 3, complete data) are kept unchanged by the zigzag algorithm and correspond to the lowest and highest mean yield of all the blocks.

AMMI preliminary analyses

Table 4 gives the ANOVA for AMMI4. The genotypes, environments and GEI account for 4.1 %, 86.4 %, and 9.5 % of the treatment sum of squares (SS). The noise in the GEI may be estimated by the interaction $df$ times the error MS, namely 40.80.
Table 3 – Dominant and number of significantly dominated genotypes for JRA, environments where the genotypes were dominant (JRA) and where the genotypes were winners (AMMI). The results are for the complete data set and the incidence rates of missing values, and based on one run (out of 100) of the simulation described above. Abbreviations for the environments: Bej1: Beja1; Bej2: Beja2; Ben1: Benavila1; Ben2: Benavila2; Evo: Évora; Elv1: Elvas1; Elv2: Elvas2; Elv3: Elvas3; Rev: Revilheira; Tav1: Tavira1; Tav2: Tavira2

| Dominant or Winner genotype | JRA | AMMI |
|-----------------------------|-----|------|
|                             | Range of dominance | Number of significantly dominated genotypes | Environments |
|                             | t test* | t test** | Scheffe* | Bonferroni* |                             |
| Complete data               |       |       |          |            |                                |
| TE9115                      | [2.21; 2.27] | 3 0 0 0 | Rev, Ben1, Evo |
| TE9008                      | [2.27; 2.80] | 2 0 0 0 | Rev |
| TE9204                      | [2.80; 3.40] | 3 1 0 1 | Ben2 |
| CELTA                       | [3.40; 8.84] | 4 2 0 2 | Elv1, Bej1, Tav1, Elv2, Tav2, Elv3, Ben2, Bej2 |
|                            |       |       |          |            |                                |
| 5 % of missing values       |       |       |          |            |                                |
| TE9115                      | [2.21; 2.38] | 2 0 0 0 | Rev |
| TE9008                      | [2.38; 2.60] | 2 0 0 0 | Rev |
| TE9204                      | [2.60; 3.48] | 3 1 0 0 | Ben1, Evo |
| CELTA                       | [3.48; 8.88] | 4 2 1 2 | Elv1, Bej1, Tav1, Elv2, Tav2, Elv3, Ben2, Bej2 |
|                            |       |       |          |            |                                |
| 10 % of missing values      |       |       |          |            |                                |
| TE9008                      | [2.22; 3.17] | 4 1 0 1 | Ben1, Rev, Evo |
| TE9204                      | [3.17; 3.64] | 5 1 1 1 | Elv2, Ben2 |
| CELTA                       | [3.64; 9.47] | 5 2 1 2 | Elv1, Bej1, Tav1, Elv2, Tav2, Elv3, Ben2, Bej2 |
|                            |       |       |          |            |                                |
| 25 % of missing values      |       |       |          |            |                                |
| TE9115                      | [2.09; 2.10] | 5 3 0 2 | Ben1, Rev, Evo |
| TE9008                      | [2.10; 3.17] | 5 3 0 2 | Ben1, Rev, Evo |
| TE9204                      | [3.17; 3.75] | 6 4 2 3 | Elv2, Elv3, Ben2 |
| CELTA                       | [3.75; 8.77] | 6 5 2 4 | Elv1, Bej1, Tav1, Elv2, Tav2, Ben2, Bej2 |
|                            |       |       |          |            |                                |
| 50 % of missing values      |       |       |          |            |                                |
| TE9115                      | [2.07; 2.09] | 5 4 1 2 | Ben1, Rev, Evo |
| TE9008                      | [2.09; 3.16] | 2 2 1 2 | Ben1, Rev, Evo |
| TE9204                      | [3.16; 3.85] | 3 3 2 3 | Elv2, Elv3, Ben2 |
| CELTA                       | [3.85; 9.21] | 3 3 2 3 | Elv1, Bej1, Tav1, Tav2, Bej2 |
|                            |       |       |          |            |                                |
| 75 % of missing values      |       |       |          |            |                                |
| TE9115                      | [1.52; 3.47] | 8 8 8 8 | Ben1, Rev, Eov, Ben2 |
| TE9204                      | [3.47; 9.10] | 8 8 8 8 | Ben1, Rev, Eov, Elv1, Bej1, Tav1, Elv2, Tav2, Elv3, Ben2, Bej2 |

*0.05; **0.01

which by difference from the total of 141.74 (total GEI SS) implies a GEI signal SS of 100.94, or 71.21 % (Gauch, 1992). Figure 2 shows the numbers of indirect replications for the AMMI model family from AMMI0 to AMMI8. The models are less parsimonious, or more complex, moving to the right. AMMI2 achieves the highest number of indirect replications of 1.66 (i.e. 1 replication gives 1.66 more information when considering the parsimonious AMMI2 model). To the left of this model, excessively simple models underfit the real signal, whereas to the right, excessively complex models overfit the spurious noise. This relationship between accuracy and parsimony has been named as Ockham’s hill (Gauch, 2006; Mackay, 1992).

Since the signal is much simpler than the noise, the signal is extracted selectively in early model parameters whereas noise is extracted selectively in late model parameters. A parsimonious model, which captures most signals and discards most of the noise, can be chosen by stopping at the right point (Gauch, 1992). From Table 4 it is possible to obtain the SS of the GEI signal of 100.94 (“total GEI SS” minus “noise in GEI”) and the SS for the first two PCs together of 115.05 (77.04 for IPC1...
and 38.01 for IPC2), which means that these two PCs are mostly signal whereas the remaining are mostly noise. The F tests in Table 4 also suggested retaining the first two PCs. For comparison with AMMI, the Finlay-Wilkinson linear regressions on environment mean capture a SS of 43.63, which is about 56.6 % of the GEI SS captured by IPC1.

Figure 3 depicts the AMMI1 biplot for the durum wheat experiment. The choice of the AMMI1 biplot instead of AMMI2 was made to allow the comparison with Figure 1. The absissa shows the main effects and the ordinate shows the IPC1 scores. The 9 genotypes are represented in bold font and the 11 environments (Bej1: Beja1; Bej2: Beja2; Ben1: Benavila1; Ben2: Benavila2; Rev: Revilheira; Elv1: Elvas1; Elv2: Elvas2; Elv3: Elvas3; Rev: Revilheira; Tav1: Tavira1; Tav2: Tavira2) which are dominated by the values until 50 %. While for JRA there are six environments (Rev, Ben1, Tav1, Tav2 and Bej2) which are dominated by the same genotypes decreases when increasing the proportion of missing values. The only exception is the case with 75 % of missing values, with 6 agreements between analyses, which is more likely to change each time the random procedure to remove observations is run.

Regarding the comparison between percentages of missing values, Table 3 (second, eighth and ninth columns) illustrates a more stable and robust performance of JRA, since the dominant genotypes are kept unchanged for an incidence of missing values until 50 %. While for JRA there are six environments (Rev, Elv1, Bej1, Tav1, Tav2 and Bej2) which are dominated by the

The analysis of Table 3 should be performed between methods and between incidence rates of missing values. Regarding the comparison between methods, the most similar results are for the complete data without missing values, with eight environments having higher yield for the same (dominant/winner) genotypes. The number of environments dominated/won by the same genotypes decreases when increasing the proportion of missing values. The only exception is the case with 75 % of missing values, with 6 agreements between analyses, which is more likely to change each time the random procedure to remove observations is run.

Stability with missing values

JRA is an extremely robust technique against missing observations in what concerns genotype comparison and selection (Pereira et al., 2007). They used a series of 17 experiments of x–designs of winter rye genotypes, in the years of 1997 and 1998, and considered proportions of missing values from 5 % to 75 %, with step size of 5 % generated randomly in triplicate. The durum wheat data set was used here to test the stability and agreement in choosing the dominant genotypes for different incidence rates of missing values, between JRA and AMMI. Table 3 presents the main results for different incidence rates of missing values. The missing values were chosen randomly as described before.

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Figure 3 – AMMI1 biplot for the durum wheat experiment. Bold font represents the codes of the genotypes and plain text the abbreviations for the environments (Bej1: Beja1; Bej2: Beja2; Ben1: Benavila1; Ben2: Benavila2; Evo: Evora; Elv1: Elvas1; Elv2: Elvas2; Elv3: Elvas3; Rev: Revilheira; Tav1: Tavira1; Tav2: Tavira2).
same genotypes in all the cases (with exception of the extreme 75% incidence rate of missing values), for the AMMI analysis it only happens in 4 environments (all of them are won by CELTA). Moreover for the AMMI model the genotype TE9008 and TE9115 only win in one of the five cases (incidence rates), while for the JRA the dominant genotypes are more stable.

Although the dominant genotypes have little change with the incidence rate of missing values it seems clear that CELTA is the strongest genotype regarding the yield production. It is always dominant for higher environmental indexes and always wins one mega-environment. With 75% of missing values (297 out of 396 observations) the JRA yet identifies two of the dominant genotypes presented in the upper contour of Figure 1, while AMMI identifies a “small” mega-environment Elv3 and a larger mega-environment with the remaining ten environments (Table 3).

We carried out 100 simulations as described before, and Table 3 shows the results for one of them chosen randomly. The 100 data sets for each proportion of missing values resulted in the identification of, at least, one dominant/winner genotype coincident to the complete data set when considering 75% of missing values. For 50% or less JRA always identified TE9008 and CELTA as dominant genotypes, whereas TE9204 (not dominant/winner in the complete data set) and CELTA almost always win one AMMI mega-environment. A detailed summary of the 100 runs is presented in Table 5.

Table 5 – Proportion of runs in which dominant genotypes (JRA) and winners of mega-environments (AMMI) are common to the results of the original data.

| Proportion of missing values | Dominant or Winner genotype | JRA | AMMI |
|-----------------------------|-------------------------------|-----|------|
| 5%                          | TE9115                        | 7   | 28   |
|                             | TE9008                        | 100 | 47   |
|                             | TE9204                        | 78  | 93   |
|                             | CELTA                         | 100 | 100  |
| 10%                         | TE9115                        | 8   | 14   |
|                             | TE9008                        | 100 | 71   |
|                             | TE9204                        | 56  | 98   |
|                             | CELTA                         | 100 | 100  |
| 25%                         | TE9115                        | 12  | 9    |
|                             | TE9008                        | 100 | 62   |
|                             | TE9204                        | 72  | 100  |
|                             | CELTA                         | 100 | 100  |
| 50%                         | TE9115                        | 21  | 15   |
|                             | TE9008                        | 100 | 36   |
|                             | TE9204                        | 43  | 94   |
|                             | CELTA                         | 100 | 100  |
| 75%                         | TE9115                        | 3   | 34   |
|                             | TE9008                        | 19  | 29   |
|                             | TE9204                        | 84  | 41   |
|                             | CELTA                         | 98  | 100  |

**Conclusion**

The aim was not to compute estimates of missing values and compare them with the original data, but to compare the final results (i.e. dominant/winner genotypes and environments where the were dominant/winner) between JRA and AMMI and between the complete data and incomplete data sets with different incidence rates of missing values. The main conclusions were the similarity between the dominant genotypes in JRA and the winners of the mega-environments in the AMMI analysis; and a more stable performance of JRA for higher proportions of missing values. The results from JRA trend to be more significant than those from AMMI models in these kind of trials, because the genotypes in the program have proved to have strong adaptability.

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