Artificial neural networks compared with Bayesian generalized linear regression for leaf rust resistance prediction in Arabica coffee

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Abstract – The objective of this work was to evaluate the use of artificial neural networks in comparison with Bayesian generalized linear regression to predict leaf rust resistance in Arabica coffee (Coffea arabica). This study used 245 individuals of a F2 population derived from the self-fertilization of the F1 H511-1 hybrid, resulting from a crossing between the susceptible cultivar Catuaí Amarelo IAC 64 (UFV 2148-57) and the resistant parent Híbrido de Timor (UFV 443-03). The 245 individuals were genotyped with 137 markers. Artificial neural networks and Bayesian generalized linear regression analyses were performed. The artificial neural networks were able to identify four important markers belonging to linkage groups that have been recently mapped, while the Bayesian generalized model identified only two markers belonging to these groups. Lower prediction error rates (1.60%) were observed for predicting leaf rust resistance in Arabica coffee when artificial neural networks were used instead of Bayesian generalized linear regression (2.4%). The results showed that artificial neural networks are a promising approach for predicting leaf rust resistance in Arabica coffee.

Index terms: Coffea arabica, Hemileia vastatrix, artificial intelligence, molecular markers, prediction.

Introduction

Coffee is a culture that belongs to the Coffea genus of the Rubiaceae family. It is one of the main Brazilian crops, placing Brazil as the largest coffee producer and exporter in the world, and as the second largest coffee consumer. Among the cultivated species, Coffea arabica L. stands out for presenting great economic value, since it is the most appreciated coffee by consumers (Brasil, 2015). Factors that could affect coffee production include major diseases, such as the coffee leaf rust or coffee berry disease (Silva et al.,...
2006). Particularly, leaf rust is the main disease that affects coffee, and it is caused by *Hemileia vastatrix*, which affects all coffee producing regions worldwide (Várzea et al., 2002). One way to minimize the damage caused by this disease is to use resistant varieties, obtained in breeding programs (Alvarenga et al., 2011). In this context, plant genome sequencing and identification of molecular markers associated with resistance have been used to assist breeding programs (Morceli et al., 2008; Alvarenga et al., 2011).

Among the statistical methods based on molecular markers, genome wide selection (GWS) is a technique that has been widely used for selecting materials with higher performance based on information from molecular markers. Such methodology, proposed by Meuwissen et al. (2001), allows incorporating molecular information directly in the prediction of the individuals’ genetic merit. In general, the methodologies developed for genomic selection, such as Ridge Regression Best Linear Unbiased Prediction (RR-BLUP), Bayes A and B (Meuwissen et al., 2001) and Bayesian Lasso (Campos et al., 2009), are based on the normality assumption of the studied phenotype. Thus, using them in categorical variables, such as leaf rust resistance, becomes inadequate.

In order to avoid this limitation, Pérez & Campos (2014) proposed the use of Bayesian generalized linear regression (BGLR), extending the genomic selection to continuous (e.g., yield) and discrete (e.g., disease resistance) models (Pérez & Campos, 2014). However, even including other types of models, the methodology requires that these models are previously established by the researcher, i.e., the problem’s modeling is essential to the use of this technique. Also, another deficiency of the techniques commonly used in GWS is that they do not extend to problems involving dominance and epistatic effects (Resende, 2007).

As an alternative to the use of conventional stochastic modeling, artificial neural networks can be used for prediction and classification, with the advantage of not having assumptions regarding the model that should be adopted, since their results depend on the learning process rather than on the distribution of variables. Such approach has been successfully applied to solve several problems related to genetics (Nascimento et al., 2013; Silva et al., 2014, 2016; Sant’Anna et al., 2015).

The objective of this work was to evaluate the use of artificial neural networks in comparison with Bayesian generalized linear regression to predict leaf rust resistance in Arabica coffee (*Coffea arabica*).

### Material and Methods

Phenotyping and genotyping experiments were carried out in a greenhouse and in a coffee biotechnology laboratory, the Laboratório de Biotecnologia do Café (BioCafé), of the Instituto de Biotecnologia Aplicada à Agropecuária (Bioagro), an institute of biotechnology applied to agriculture, of the Universidade Federal de Viçosa (UFV) (20°45’37”S, 42°52’4” W) as described by Pestana et al. (2015). Phenotyping was conducted in May 2009, July 2009 and August 2009 for the repetitions 1, 2 and 3 respectively. Genotyping was carried out in the years 2010, 2011 and 2012. The study used 245 individuals in a F2 population derived from the self-fertilization of the F1 H511-1 hybrid, resulting from a crossing between the susceptible cultivar Catuaí Amarelo IAC 64 (UFV 2148-57) and the resistant parent Hibrido de Timor (UFV 443-03). DNA extraction of F2 population individuals was carried out according to the protocol described by Diniz et al. (2005). Genotyping was carried out with 137 markers (74 AFLP, 58 SSR, 4 RAPD, and 1 specific primer) (Pestana et al., 2015).

Markers data for each individual were scored for genomic selection analysis. For approaching dominant markers (allele from the resistant parent Hibrido de Timor UFV443-03), -1 was assigned for presence, and 1 for absence of the band. For repulsion dominant markers (allele from the susceptible parent Catuaí Amarelo, UFV 2148-57), 1 and -1 were assigned for presence and absence of the band, respectively. The codominant markers were scored as 0 for heterozygote; -1, for bands from the resistant parent; and 1, for bands from the susceptible parent.

In the phenotyping regarding the expression of genotype resistance or susceptibility, inoculations were carried out, and uredospores of *H. vastatrix* race II were multiplied and inoculated according to the methodology described by Capucho et al. (2009).

Inoculation – for the 245 individuals of a F2 – was carried out using leaf discs, as described by Eskes (1982), using 20 µL of spore suspension with viability over 30% at a concentration of 2.0 mg µL-1, with three replications. In each plant and each replication, 16 leaf discs of 1.5 cm diameter were used as sample units,
and four discs from each plant were uninoculated as a control. After inoculation, the discs were placed on a nylon fabric with water and saturated foam inside a gearbox disinfected, followed by methodology described by Capucho et al. (2009).

The assessment of resistance/susceptibility was carried out according to Capucho et al. (2009), following the scale of Tamayo et al. (1995), which is based on the absence or presence of uredospores. According to this scale, scores 1-3 are considered resistant (1 - absence of symptoms; 2 - small chlorotic lesions; and 3 - large chlorotic lesions without sporulation). Susceptibility (scores 4 to 6) is assigned to large chlorotic lesions, with few uredospores, occupying less than 25% of the leaf area (score 4); to lesions with sporulation, occupying 25 to 50% of the leaf area (score 5); and to lesions with sporulation, occupying more than 50% of the leaf area, and with uredospores (score 6). To predict the resistance pattern to leaf rust using two methodologies (BGLR and ANN), scores were rescored as resistant (1 = scores 1 to 3) and susceptible (0 = scores 4 to 6).

The architecture of the ANN used was the single hidden layer back-propagation (Rumelhart et al., 1986) with six neurons, 137 inputs (represented by the marker values) and one output layer, which provided the prediction of leaf rust resistance. The architecture was represented by a functional form, as the topology described in Figure 1.

The $W_m$ variables were functions of pondered sums of the 137 input variables $X_i$ (represented by the marker values), and the output variable, $Y_k$ (leaf resistance scores), that is, $W_m = \gamma(\alpha_m + \alpha_m^B X_i)$, $m = 1, 2, ..., 6$, which was modeled as functions of these combinations.

$T_i = \beta_k + \beta_k X_i W_m$; $k = 1, 2, ..., K$;

$Y_k = f_k(X_i) = g_k(T_i), k = 1, 2, ..., K$;

in which $W = (W_1, W_2, ..., W_m)$ and $T = (T_1, T_2, ..., T_k)$. The activation function $\gamma(\cdot)$ and the output function $g_k(T)$ were, respectively, the sigmoid, $\gamma(\cdot) = 1/(1+e^{-\theta})$, and the softmax, $g_k(T) = e^T / \sum_{i=1}^k T_i$ (Hastie et al., 2009).

The estimate of the weights $\{\alpha_m, \alpha_m^B; m = 1, 2, ..., M\}$ and $\{\beta, \beta_k; k = 1, 2, ..., K\}$ was carried out by minimization of the sum of square errors,

$$R(\theta) = \sum_{k=1}^{K} \sum_{i=1}^{N} (y_k - f_k(x_i))^2,$$

by the application of the descending gradient algorithm. Since the hidden layers were used, the adjustment procedure known as back-propagation was applied.

For the network training (obtainment of weights), the set of observations was divided in two parts: training and validation, as commonly adopted in the literature (Silva et al., 2014, 2016; Sant’Anna et al., 2015). The first, denoted by training set, consisted of 160 individuals taken at random. The second, composed of the remaining 85 observations, was used for network validation. The number of individuals for training and validating the network was defined following the percentages (around 65% of the data information to train the network) usually adopted by the other authors (Braga et al., 2011; Nascimento et al., 2013). The arguments required for the function of the network, such as number of neurons in the hidden layer, weight initial value, and decay rate were chosen considering the network that provided error value of 15%, at most, for the validation set.

For comparison, prediction was also carried out using the genomic selection method based on generalized linear models under the Bayesian approach (Campos & Perez Rodriguez, 2014). Regression model under the Bayesian approach is given by: $y_i = \eta_i + \epsilon_i$, where $\eta_i$ is a

![Figure 1. Schematic of a single hidden layer back-propagation.](image)
linear predictor, and $\epsilon_i$ are independent normal model residuals.

Rewriting $\eta_i$: $\eta_i = \mu + X_1 \beta_1 + \ldots + X_j \beta_j + u_1 + \ldots + u_q$, in which $\mu$ is an intercept; $X_j$ are the matrices for predictors, $X_p = \{x_{ijk}\}$; $\beta_j$ are the vectors of effects associated to the columns of $X_j$; $u_q = \{u_{q1}, \ldots, u_{qn}\}$ are the vectors of random effects.

For categorical variables, as leaf rust resistance, the vector response adopted in BGLR is the probit link (Perez & Campos, 2014), where the probability of each of the categories is linked to the linear predictor according to the link function below: $P(y_i = k) = \Phi(\eta_i - y_k) - \Phi(\eta_i - y_{k-1})$, where $\Phi(.)$ is the standard normal cumulative distribution function, $\eta_i$ is the linear predictor, and $y_k$ are threshold parameters ($y_0 = -\infty$, $y_k \geq y_{k-1}$ and $y_k = \infty$).

The prior density was the same adopted in the Bayesian Lasso, which assumes that the marginal distribution of marker effects is double exponential (Perez & Campos, 2014).

To estimate the model in BGLR, the study used the same 65% of individuals used in ANN training. For validation, the study used the same 35% of individuals used to validate the neural network.

After obtaining the results, the percentage of prediction errors was calculated for the artificial neural networks and the Bayesian generalized linear regression was studied in training populations (estimate) and in validation. Coincidence coefficients between predictions were calculated regarding leaf rust resistance by both approaches. Manhattan plots were constructed in order to observe if the behavior and distribution of the most important markers for neural networks and Bayesian approach were similar.

In order to verify the possibility of using marker assisted selection for predicting leaf rust resistance, two situations were taken into account. At first, the study considered the three most important markers, for which three prediction scenarios were assessed: the use of the most important marker; the use of the two most important markers; and the use of the three most important markers.

In the second assessed situation, among the ten most important markers, the study selected the two markers that belong to the linkage groups in which Pestana et al. (2015) detected two important QTLs for leaf rust resistance in *C. arabica*.

The models used were carried out by using nnet (to fit a neural network) function and BGLR (to fit a generalized Bayesian model) functions of the packages nnet (Venables & Ripley, 2002) and BGLR (Campos et al., 2009; Pérez et al., 2010), respectively. Both were developed in the R software (R Core Team, 2016).

A total of 100,000 iterations were used in this study, with 20,000 burn-in and thin iterations assuming the value 10, following authors as Macpherson et al. (2004) and Kwon & Reis (2015).

### Results and Discussion

After the training algorithm was executed, the topology with prediction error rate of less than 15% was considered. For this topology, the number of neurons was six, decay rate was $5.0 \times 10^{-4}$, and maximum number of iterations was equal to 5,000. The initial values for the network weights were randomly chosen from the interval $[-0.5, 0.5]$ and assuming max = 2. According to Venables & Ripley (2002), the initial values of the process should be chosen in order to satisfy the equation $LS \times \max(|x|) \approx 1$, where LS denotes the upper limit of the range and $\max(|x|)$ is the largest absolute value of the set of training data.

Neural networks provided slightly lower error rates than BGLR in the validation population. Specifically, error rates were equal to 2.40% and 1.60% in the validation population, considering the predictions carried out by BGLR and ANN, respectively (Table 1).

| Methodology                                 | Classification error (%) |
|---------------------------------------------|--------------------------|
| Artificial neural network                   | Training: 0.00, Validation: 1.600 |
| Bayesian generalized linear regression (BGLR) | Training: 0.00, Validation: 2.400 |
| Concordance (%)                             | - 81.37                  |

Results and Discussion

In order to select resistant varieties, genomic selection methodologies, especially Bayesian generalized linear regression, once that BGLR extend the genomic selection to discrete variables as resistance, have been proposed to predict the breeding values by analyzing their phenotypes and high-density marker information (Heffner et al., 2009; Pérez & Campos, 2014). However, when carrying out prediction studies of genetic values
from phenotypic values with aggregation of molecular information, some statistical and genetic aspects should be considered.

In the statistical context, appropriate models must be adopted upon the possibility of occurrence of serious problems of multicollinearity and dimensionality of the incidence matrix (Ferrari, 1989). Multicollinearity is conditioned by the existence of correlations between markers and can result in inconsistent estimates of the regression coefficient and also in an overestimation of the direct effects of the explanatory variables on the dependent variable, which may lead to misinterpretation, and dimensionality issue occurs when the number of available observations is lower than the number of explanatory variables (or markers) included in the model (Cruz & Carneiro, 2003; Resende, 2007).

The existence of multicollinearity and the dimensionality issue are limiting factors for the Bayesian models, since the once that data set used in this methodology needs to be defined and known priorly, satisfying the presupposition of being in the exponential family (Resende, 2007; Garcia et al., 2012). In comparison to this, the ANN is not affect by those issues because it does not involve stochastic modeling, since it is nonparametric and the theory behind them is quite different from the linear model BGLR described above, based on computational intelligence and principles of learning (Heslot et al., 2012; Silva et al., 2014).

Apparently, both BGLR and ANN approaches are able to circumvent such problems, since results regarding prediction were satisfactory. In this study, dimensionality is not a limiting factor in view of the number of observations and studied markers. In the genetic context, it should be taken into account that BGLR approach uses a linear model in which the predicted value – being function only of favorable allele dose – predicts the additive genetic value with higher mean squared error, or lower genotypic reliability, expressed by the $r^2$, of the model because of the effects assigned to dominance and epistatic interactions disregarded in this model (Cruz, 2012). However, when a computational intelligence approach as ANN is used, such information and other important information that the breeder has about the genotype in study constitute a relevant input to be considered during the training processes for increasing the efficiency of the selection process, and it becomes possible to capture different relationships between markers and phenotypes (Bureau et al., 2005; Heslot et al., 2012; Silva et al., 2014, 2016).

The percentage of agreement between the results obtained by the two methodologies was 81.37%. Comparing the results obtained with the mapping that had recently been carried out by Pestana et al. (2015), neural networks were able to identify four markers (E-CGT/M-ATC1, E-CGT/M-TCT1, E-CTT/M-TGC3, and E-CGA/M-ACA2) belonging to the identified QTLs. BGLR could only identify two markers belonging to QTLs. It is believed that the ANN approach, due to its different neurons in hidden layers, can capture linear and nonlinear relationships between the response variable and the values assigned to molecular markers (Braga et al., 2011). This means that they were capable of detecting the effects of dominance and epistasis that are neglected by other methodologies (Heslot et al., 2012; Silva et al., 2014, 2016).

Manhattan plot graphs showed the most important markers for both methodologies (Figure 2). Markers behavior pattern regarding its importance in the prediction process was different in the two methodologies (ANN and BGLR) regarding leaf rust resistance. This difference occurs since the markers that were identified as the most important are different in the two methodologies. Particularly for neural networks, it is noted that a relatively large number of markers had effects of great magnitude.

The 10 most important markers obtained for the Bayesian methodology were: M34, M36, M132, M101, M88, M62, M29, M85, M65 and M116; and the 10 most important markers obtained for the neural networks were: M6, M19, M35, M44, M56, M80, M59, M64, M67 and M84. These results justify the difference between the graphs.

In order to simulate a selection strategy similar to that used in marker assisted selection (MAS) – in which only a small group of markers, supposedly in linkage disequilibrium with QTL, could be effective in prediction –, classification error rates were obtained for ANN, by evaluating three situations (Table 2). Error rates were relatively high, and ranged between 44.90% and 53.47%, evidencing the importance of using all the information available for predicting
leaf rust resistance in coffee, i.e., the need for more parameterized predictive models.

The high prediction error rates obtained in Table 2 when using markers selection show that the joint action of several markers is more appropriate for prediction of resistance or susceptibility to *H. vastatrix*. The complexity of the genetic control of *H. vastatrix* has been reported by several authors (Bettencourt & Carvalho, 1968; Capucho et al., 2009; Pestana et al., 2015). Studies on coffee demonstrated that Hibrido de Timor derived accessions have five dominant genes (SH5, SH6, SH7, SH8 and SH9) that provide, alone or in combination, the resistance spectra (Bettencourt et al., 1992), indicating both horizontal control, of polygenic nature, and vertical control, provided by few genes of major effect. In this study, the most relevant markers are those that present gametic phase disequilibrium with the probable genes that control the studied trait. This association, either by factorial linkage, or by a non-random combination between marker and QTL (quantitative trait loci), has been considered a crucial factor in obtaining gains with selection (Resende et al., 2008).

In general, important markers to be used in MAS are identified through genomic analyses in populations structured for genetic mapping with the aid of regression models, assuming the existence of markers flanked to significant QTLs (Caixeta et al., 2009). In this work, the use of computational intelligence as an auxiliary tool for QTLs detection was proposed. For assisted selection of this work, the ten most important markers for prediction were selected. Another reason for the high prediction error rates found in Table 2, even for the markers identified in the linkage maps, is the fact that the QTLs identified by Pestana et al. (2015) have low coefficients of determination (5.8% and 8.1%). This means that they are not sufficient to confer leaf rust resistance in *C. arabica* alone.

Finally, it is noteworthy that the low polymorphism rate found in *C. arabica* populations (Geleta et al., 2012) was not a limiting factor for the use of neural networks in predicting leaf rust resistance, since it presented satisfactory results considering the prediction error rate. The proposed method has low prediction error rate and superior results when compared to those found by the Bayesian generalized model. The advantage of neural networks over other approaches has been proven in genetic studies. In relation to genotype x environment interaction, Nascimento et al. (2013) assessed the adaptability and phenotypic stability of alfalfa genotypes based on formation of an artificial

**Figure 2.** Importance of markers for the prediction process: A, graph in which the x axis corresponds to the information of the markers, and the y axis represents the estimate of markers effect for the model obtained by Bayesian generalized linear regression; and B, graph in which the x axis represents the dispersion of markers; and the y axis represents the respective prediction errors when each marker is removed from the model obtained by artificial neural network.

|        | Situation I | Situation II |
|--------|-------------|-------------|
| MAS (1 marker) | TEP | MAS (2 markers) | TEP | MAS (3 markers) | TEP | MAS (3) TEP |
| -1 | 51.02 | (1,1) | 44.9 | (1,1,1) | 43.67 | (1,1) | 46.12 |
| -1 | 48.98 | (1,1) | 46.94 | (1,1,1) | 47.35 | (1,1) | 46.94 |
| -1 | 52.24 | (1,1) | 53.47 | (1,1,1) | 53.88 | (1,1) | 47.75 |
| -1 | - | (1,1) | 48.16 | (1,1,1) | 44.9 | (1,1) | 46.53 |
| -1 | - | (1,1) | 46.12 | (1,1) | 46.12 | (1,1) | 46.12 |
| -1 | - | (1,1) | 46.94 | (1,1) | 46.94 | (1,1) | 46.94 |
| -1 | - | (1,1) | 47.75 | (1,1) | 47.75 | (1,1) | 47.75 |

*(Mark assisted selection by Pestana et al. (2015)).
neural network, considering the method of Eberhart & Russell (1966). Percentage agreements were of 89 and 100% for adaptability, and of 78 and 100% for stability, considering favorable and unfavorable environments, respectively. Silva et al. (2014, 2016) concluded that ANNs are efficient in predicting values and genetic gain in simulated trials under randomized block design. Regarding the classification studies, Sant’Anna et al. (2015) showed that neural networks had results superior to those obtained by discriminant analysis in the classification of simulated populations.

Conclusions

1. Artificial neural network is more efficient to predict leaf resistance in Coffea arabica than Bayesian generalized linear regression.

2. Artificial neural networks are an interesting and promising approach to predict C. arabica leaf rust resistance.

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