Associating Molecular Markers with Virus Resistance to Classify Sweetpotato Genotypes

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ABSTRACT. Molecular markers linked to resistance to sweetpotato chlorotic stunt closterovirus [SPCSV (genus Crinivirus, family Closteroviridae)] and sweetpotato feathery mottle virus [SPFMV (genus Potyvirus, family Potyviridae)] were selected using quantitative trait loci (QTL) analysis, discriminant analysis and logistic regression. Eighty-seven F1 sweetpotato [Ipomoea batatas (L.) Lam.] genotypes from a cross of ‘Tanzania’ and ‘Wagaholige’ landraces were used to generate DNA marker profiles for this study. Forty-five of the clones were resistant to SPCSV while 37 were resistant to SPFMV. A combination of 232 amplified fragment length polymorphism (AFLP) markers and 37 random amplified polymorphic DNA (RAPD) markers obtained were analyzed to determine the most informative markers. All three statistical procedures revealed that AFLP marker e41m33.a contributed the greatest variation in SPCSV resistance and RAPD marker S13.1130 accounted for most of the variation in SPFMV resistance. The power of discriminant and logistic analyses is that you do not need a parent-progeny population. An evaluation of these two models indicated a classification and prediction accuracy rates of 96% with as few as four markers in a model. Both multivariate techniques identified one important discriminatory marker (e44m41.j) for SPCSV and two markers (e41m37.a and e44m36.d) for SPFMV that were not identified by QTL analysis.

The most widely used approach to germplasm improvement in sweetpotato is mass selection (Jones, 1986). This fundamentally simple technique lends itself well to an allohexaploid crop that possesses few, if any, simply inherited traits (Buteler et al., 1999; Jones, 1986; Zhang et al., 2001). Unfortunately, self-incompatibility prevents quick introgression of desirable traits from landraces or exotics into adapted material. The preponderance of quantitatively inherited traits in sweetpotato (Jones, 1986), environmentally unstable trait expression, and the encumbrance of 2-year breeding cycles underlie recent attempts to use trait-linked DNA molecular markers in breeding programs.

The most compelling reason to develop trait-linked DNA markers in sweetpotato is to effectively breed for virus resistance. Viral diseases have been identified to cause up to 56% to 98% yield reduction in sweetpotato (Gibson et al., 1997; Ngeve, 1990). According to Geddes (1990) the sweetpotato virus disease (SPVD), a result of the co-infection of SPCSV and SPFMV, is the most destructive of the viral diseases within the African continent (Gibson et al., 1997; Karyeja et al., 1998). Hahn et al. (1981) and Mwanga et al. (2002a) estimated high broad-sense heritability for resistance to SPVD, suggesting that breeding for resistance is possible.

Ukoskit and Thompson (1997) and later Kriegner et al. (2003) demonstrated the feasibility of constructing a genetic linkage map for sweetpotato. Mapping was based on the single-dose restriction fragment (SDRF) method described by Wu et al. (1992). Progeny segregating in a 1 present :1 absent marker ratio arise from a simplex × homozygous recessive test cross (e.g., aaaaa × Aaaaaa). Progeny segregating in 4:1 ratio that generate duplex markers AAAAA are also used. A map by Kriegner et al. (2003) contained 1067 AFLP markers ordered in ~90 linkage groups with genome coverage suitable for detection of QTL. Mwanga et al. (2002a) used linkage maps to associate DNA RAPD and AFLP markers to SPFMV and SPCSV resistance. Attempts to identify markers associated with SPVD were unsuccessful because few progeny in the defined population were resistant. Despite such efforts, QTL analysis is still a challenge in breeding programs for a crop like the sweetpotato since a complete genetic map is still unavailable.

Faced with the complications associated with QTL analysis, this study seeks to determine whether the results of molecular marker and phenotype association for genotype characterization and disease resistance prediction using multivariate statistical procedures like discriminant analysis and logistic regression are comparable to those of QTL analysis. Both permit assembly of test populations (i.e., resistant and susceptible classes, without regard to genetic relationships).

Discriminant analysis is a multivariate technique that uses multiple predictor variables to associate an individual with a descriptive class (Fisher, 1936). In our genetic studies of sweetpotato, discriminant analysis would involve relating a genotype,
as described by its molecular marker profile, with its disease resistance class or phenotypic group. Using the markers as the multiple variables, the method would develop an optimum classification function that maximizes differences between the resistant and susceptible classes while minimizing differences among members of those two classes. The efficiency of the function is evaluated by cross-validation as described by Lachenbruch and Mickey (1968). Cross-validation is a leave-one-out method that aims to reduce the optimistic bias in calculating classification error-rate. If the assumption of multivariate normality is not met or when no assumptions are valid about the distribution within each phenotypic group, a nonparametric procedure like k-nearest-neighbor (Rosenblatt, 1956) is used for classification. With the k-nearest-neighbor method, a Mahalanobis distance is calculated between group means and this distance is then used to determine an individual’s closeness to the mean. All n observations in the dataset are used to calculate distances; however, the observation to be classified is left out from the k-nearest-neighbors of that observation during function development.

Recent work on genetic analysis using discriminant analysis include that of Capdevielle et al. (2000), who associated microsattellites and agronomic traits in rice. Fahima et al. (2002), who investigated microsatellite polymorphism in wheat, also used discriminant analysis in their characterization.

Logistic regression is a statistical technique that is frequently used to associate explanatory variables with a binary outcome (Osir and Uchida, 2000). When there are more than two response classes, discriminant analysis is more reliable than logistic regression. Models developed using logistic regression have the advantage of summarizing and interpreting the data and hence the researcher can make inferences on factors being investigated. In our studies, the process involved relating molecular markers as the independent factors and presence or absence of viral infection as the dichotomous response. To our knowledge there is no work on logistic regression for molecular marker selection in sweetpotato. However, field disease prediction studies have extensively used logistic regression tools. Mukasa et al. (2003) used logistic regression to determine the odds ratios to detect single- or multiple-virus infections in sweetpotato genotypes.

The specific objectives of this study were to compare marker–trait association using QTL analysis, discriminant analysis, and logistic regression for DNA markers derived from a population of sweetpotato genotypes and to develop and validate genotype classification and resistance class prediction models for SPCSV and SPFMV.

Materials and Methods

Molecular marker detection and phenotypic data acquisition for the population was previously described by Mwanga et al. (2002b). The data consisted of an array of molecular marker and phenotypic information for 87 F1 genotypes derived from a cross between ‘Tanzania’ and ‘Wagabolige’ cultivars. Both ‘Tanzania’ and ‘Wagabolige’ are resistant to SPVD, although ‘Wagabolige’ is more resistant to SPVD than ‘Tanzania’ under field SPVD inoculum pressure (Mwanga et al., 2002b). Detection and confirmation of virus presence or absence in the genotypes involved using triple-antibody sandwich enzyme-linked immunosorbent assay (TAS-ELISA) test for SPCSV and nitrocellulose membrane enzyme-linked immunosorbent assay (NCM-ELISA) test for SPFMV (International Potato Center, 1990). Resistant genotypes tested negative while susceptible genotypes tested positive for virus presence (Mwanga et al., 2002b). ‘Tanzania’ is widely grown in sub-Saharan Africa while ‘Wagabolige’ is more confined to Uganda. The 87 F1 genotypes had 269 dominant molecular markers, 232 of them being AFLP fragments and 37 being RAPD fragments. These were two markers less than what Mwanga et al. (2002b) used. Fifty genotypes were classified as resistant and 37 susceptible to SPFMV. Forty-two were classified as resistant and 45 susceptible to SPCSV, respectively. Phenotypic data for SPCSV and SPFMV were obtained using ELISA tests and were coded as a one (1) for positive virus infection or zero (0) for negative results.

Statistical analyses were done on SPFMV and SPCSV only because only three out of 87 genotypes were resistant to SPVD. These were too few for the statistical procedures to converge and give a solution.

QTL analysis was done using QGENE (Nelson, 1997). Single-point analysis of variance (ANOVA), or regression on a marker (Soller et al., 1976), was conducted to reveal significant (P<0.05) associations between markers and virus resistance. Markers were considered independent variables and virus resistance score as the dependent variable. Proportion of the total phenotypic variance accounted for by the markers was estimated by the coefficient of determination (R2) obtained from simple linear regression (Nelson, 1997; Soller et al., 1976).

Step discriminant analysis using the STEPDISC procedure (SAS, 2001) was used to select markers to be included in the classification model. As described in SAS (1999), the forward selection process commences with no markers in the model. Entry-significant levels of P ≤ 0.03 and P ≤ 0.02 for SPCSV and SPFMV, respectively, of the chi-square score for entering an effect or marker into the model to achieve at least 95% prediction accuracy were imposed to choose the most discriminating markers (SAS, 1999). At each step the procedure enters the marker that contributes most to the discriminatory power of the model as measured by Wilks’ lambda, the likelihood ratio criterion. The process stops when all unselected markers fail to meet the entry condition. The selected markers were then used in a nonparametric discriminant analysis (k = 1), DISCRIM option (SAS, 2001), to construct and validate a class prediction function and to predict group membership, resistant or susceptible, of the test. Since the categorical nature of the data could not allow assumption of normal distribution to be made a nonparametric method, the k-nearest-neighbor method (Rosenblatt, 1956) was used to estimate the group-specific densities that produce a classification criterion.

The performance of the discriminant criterion was evaluated by group-specific error-count estimates during cross-validation. The error estimator gives the proportion of misclassified observations in each group. Total error, from which we derive percent correct classification, is the weighted mean of error estimates of the two phenotypic groups.

PROC LOGISTIC (SAS, 2001) was used to perform logistic regression with the logit link to select markers that significantly accounted for phenotypic variation, with the forward selection option used for marker selection. With forward selection, PROC LOGISTIC first estimates parameters for effects, markers in our case, forced into the model. These effects are the intercept and the first n explanatory effects in the model; n is zero by default. Next, the procedure computes the score chi-square statistic for
Results and Discussion

Sweetpotato chlorotic stunt virus. All three statistical procedures selected two common markers (e41m33.a and e38m36.u) that had a significant association with resistance to SPCSV (Tables 1, 2, and 3). These results are in concordance with those of Mwanga et al. (2002b). Marker e38m36.u was selected by discriminant and logistic but not QTL procedures as one of the informative markers. Marker e41m33.a was the most informative, regardless of the method of selection used, accounting for 66.75% of the variation in SPCSV resistance according to QTL analysis. Our results suggest the possibility of using these multivariate methods for identifying powerful QTLs. With the six markers selected by discriminant analysis to create a classification model, we achieved 96% correct classification for SPCSV. Discriminant analysis misclassified one genotype out of 42 that were phenotypically classified as being resistant. This model also misclassified into the resistant class three genotypes out of 45 that were visually described as susceptible. The model developed using logistic regression fitted the data well according to the Hosmer and Lemeshow (Hosmer et al., 1991) goodness-of-fit test ($\chi^2 = 0.082; \text{df} = 3; P = 0.9939$). Table 4 shows observed vs. expected class values for SPCSV and SPFMV.

Sweetpotato feathery mottle virus. Marker S13.1130 was identified by all three selection procedures and accounted for 64.62% of the variation in SPFMV resistance (Tables 1, 2, and 3) according to QTL analysis. Marker e39m32.f was the second most powerful marker selected by QTL analysis but neither by discriminant nor logistic analyses. Markers e41m37.a and e44m36.d were selected as being informative by discriminant and logistic analyses only (Tables 2 and 3) but in differing order of importance. The discriminant analysis model constructed using the four selected markers achieved a 96% correct classification rate. Four clones out of 50 that were phenotypically classified as being resistant were classified by discriminant analysis as being susceptible. These are the clones that discriminant analysis misclassified and they give us the total error rate of the model developed. This model achieved a highly accurate classification rate and with only four variables was a desirable result because we aim to have as few markers as possible to create an efficient class prediction model. Results of Hosmer and Lemeshow test for sweetpotato chlorotic stunt closterovirus (SPCSV) and sweetpotato feathery mottle virus (SPFMV) in sweetpotato.

Table 1. QTL analysis for DNA markers associated with ≥10% resistance to sweetpotato chlorotic stunt closterovirus (SPCSV) and sweetpotato feathery mottle virus (SPFMV) in sweetpotato.

| Disease | Marker      | F   | $R^2$ | P      |
|---------|-------------|-----|-------|--------|
| SPCSV   | e41m33.a    | 170.66 | 0.67  | <0.0001|
|         | e40m34.c    | 111.16 | 0.57  | <0.0001|
|         | e38m36.u    | 19.24  | 0.18  | <0.0001|
|         | e36m49.a    | 14.42  | 0.14  | 0.0003 |
|         | e35m49.d    | 12.58  | 0.13  | 0.0006 |
|         | R9.650      | 9.41   | 0.10  | 0.0029 |
|         | e39m40.e    | 9.23   | 0.10  | 0.0032 |
| SPFMV   | S13.1130    | 155.28 | 0.65  | <0.0001|
|         | e39m32.f    | 52.49  | 0.38  | <0.0001|
|         | e36m59.a    | 10.53  | 0.11  | 0.0017 |
|         | e33m59.a    | 9.76   | 0.10  | 0.0024 |

Table 2. GENERALISER marker selection for DNA markers associated with ≥10% resistance to sweetpotato chlorotic stunt closterovirus (SPCSV) and sweetpotato feathery mottle virus (SPFMV) in sweetpotato.

| Disease | Marker | Wilk’s lambda | P     |
|---------|--------|---------------|-------|
| SPCSV   | e41m33.a | 0.33          | <0.001|
|         | e38m36.u | 0.28          | <0.001|
|         | K2.650   | 0.26          | <0.001|
|         | AL.3.1300 | 0.24         | <0.001|
|         | e44m.41.j | 0.23         | <0.001|
|         | e39m.41f  | 0.22          | <0.001|
| SPFMV   | S13.1130 | 0.34          | <0.001|
|         | e41m37.a | 0.31          | <0.001|
|         | e40m36.d | 0.29          | <0.001|
|         | e44m36.d | 0.26          | <0.001|

Table 3. Logistic regression selection for DNA markers associated with ≥10% resistance to sweetpotato chlorotic stunt closterovirus (SPCSV) and sweetpotato feathery mottle virus (SPFMV) in sweetpotato.

| Disease | Marker selected | $\chi^2$ score | P     |
|---------|----------------|----------------|-------|
| SPCSV   | e41m33.a       | 58.49          | <0.0001|
|         | e38m36.u       | 11.36          | 0.0007 |
|         | H14.700        | 7.44           | 0.0064 |
|         | e44m41.j       | 6.61           | 0.0101 |
|         | e40m41.c       | 5.71           | 0.0169 |
|         | B7.1130        | 5.47           | 0.0194 |
|         | e32m60.g       | 5.31           | 0.0211 |
| SPFMV   | S13.1130       | 57.38          | <0.0001|
|         | e40m38.b       | 8.90           | 0.0029 |
|         | e44m36.d       | 8.56           | 0.0034 |
|         | e41m37.a       | 7.56           | 0.0060 |
|         | e39m45.c       | 6.98           | 0.0083 |
|         | e39m39.e       | 6.00           | 0.0143 |

Table 4. Partition for the Hosmer and Lemeshow test for sweetpotato chlorotic stunt closterovirus (SPCSV) and sweetpotato feathery mottle virus (SPFMV) in sweetpotato.

| Disease | Observed clones (no.) | Expected clones (no.) | Disease resistance group |
|---------|-----------------------|-----------------------|--------------------------|
| SPCSV   | 45                    | 44.98                 | 1 (present)              |
| SPCSV   | 42                    | 42.02                 | 0 (absent)               |
| SPFMV   | 37                    | 36.98                 | 1 (present)              |
| SPFMV   | 50                    | 50.02                 | 0 (absent)               |
must be performed separately for each population. In a review of QTL methods in plants, Kearsey and Farquhar (1998) showed that analytical methods locate QTL with poor precision [10–30 centimorgans (cM)], unless the heritability of an individual QTL is high. They further stated that this unreliability of QTL location may suggest false candidate genes. In his review of QTL analysis, Young (1999) states that phenotypic mis-scoring of even a few individuals can totally confound QTL discovery and placement. He also suggests that nonparametric categorical data may not be as amenable to QTL analysis as normally distributed quantitative data, thus underscoring the difficulties associated with using QTL for marker-assisted selection.

Mwanga et al. (2002b) reported studies combining both quantitative genetics and molecular marker investigations in an attempt to describe inheritance of resistance to SPCSV, SPFMV, and SPVD. However, these workers were faced with the undesirable difficulty of constructing genetic maps using null alleles and having insufficient plant numbers in the SPVD resistant category. In their QTL studies on molecular markers, Mwanga et al. (2002b) found that marker e4.1m33.a was responsible for 70% of the variation in SPCSV resistance. They also found that marker S13.1130 accounted for 72% of the resistance to SPFMV. Their conclusions were based on single-point ANOVA and multiple-regression analyses with two markers in the model. Unfortunately, a lack of progeny in the SPVD resistant class prevented disclosure of markers linked to SPVD resistance.

We attribute the slight differences between our QTL results and those of Mwanga et al. (2002b) to the fact that the data set that we used contained two markers less. Nonetheless our conclusions concur with theirs and do confirm the power of markers e4.1m33.a and S13.1130 to influence resistance to SPCSV and SPFMV, respectively.

Discriminant and logistic regression analyses have greater commonality with each other than with QTL analysis as far as markers selected is concerned possibly because the former two are both multivariate techniques. These multivariate techniques determine the power of a marker while other markers are still in the model hence giving a more accurate comparative analysis compared to the single point ANOVA for QTL analysis. Single point ANOVA is also referred to as regression on a marker and it assumes the presence of a single QTL on the genome (Broman, 2001). The ANOVA calculates the variance at marker loci without considering the effect of the other markers when they act together, hence the value obtained may not be the true effect of the QTL. In his study on genetic mapping of agronomic traits from the interspecific cross of *Oryza sativa* L. and *Oryza glaberrima* Steud., Aluko (2003) concluded that discriminant analysis was superior to interval mapping in selecting markers that could be used in grouping doubled haploid lines of rice into predefined groups.

We suggest that logistic regression is an increasingly valuable tool in molecular marker selection protocols in a sweetpotato breeding program. Investigators in the medical sciences have exploited the power of logistic regression to select important markers. In a study involving boars (*Sus scrofa* L.), Thurston et al. (2002) identified 16 candidate genetic markers (*P* ≤ 0.005) by comparing the AFLP profile with semen freezability trait using logistic regression analysis. These findings support the hypothesis that there is a genetic basis for variation in postthaw semen quality between individuals, and that AFLP technology may be able to identify molecular markers linked to genes influencing this variation. Dunsmuir et al. (2000) used logistic regression modeling to identify three immunohistochemistry markers that were statistically significant predictors of the metastatic status (M-stage, bone metastasis vs. no bone metastasis) of prostate cancer in men.

Results from our work suggest that it is possible to use logistic and discriminant multivariate techniques to select powerful markers that may be useful to breeders. While multivariate techniques may not replace QTL analysis, the flexibility of assembling test populations, regardless of relatedness, is particularly useful when progeny from a controlled cross rarely arise as resistant (e.g., SPVD progeny). Our results demonstrated that multivariate techniques can achieve results that are comparable to those of QTL analysis, even in the absence of a genetic map, and identify markers not disclosed through QTL analysis.

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