A Maximum-Type Association Test for Censored Time-to-Event Data

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Abstract

Background: Testing the association between a diallelic marker and a censored time-to-event trait is a specific problem in population-based association studies. For a certain gene, the mode of inheritance may be of particular interest. Therefore, the principle of maximum-type tests (or minimum p procedure) is modified for continuous traits, especially for censored time-to-event data.

Results: We propose a Marcus-type multiple contrast test for a single censored time-to-event trait in a population-based study assuming a Cox proportional hazard model. Using simulations we worked out the limitation of this asymptotic approach: sufficient sample sizes and non-rare alleles are required. A user-friendly implementation of this method is available in the survival and multcomp packages of the statistical software R.

Conclusions: The proposed approach can be used for the analysis of individual SNPs when censored time-to-event data in population-based association studies are of interest. The approach allows both a global claim of association and determination of the particular underlying mode of inheritance. The mode-specific hazard ratios and their lower simultaneous confidence limits provide information about statistical significance and genetic relevance.

Keywords: Max-test; Censored time-to-event trait; Population-based association study

Introduction

Particular SNPs in population-based association studies can be analyzed using a maximum test, such as the MAX-3 test. This approach is widely used for proportions in the case control design and for continuous traits. Here, an extension is proposed for censored time-to-event traits using a Marcus-type multiple contrast test under the assumptions of the Cox proportional hazard model. Simulations revealed serious limitation of this asymptotic approach as sufficient sample sizes and non-rare alleles are required. Both a global claim of association and the particular underlying mode of inheritance can be identified. The mode-specific hazard ratios and their lower simultaneous confidence limits provide information about statistical significance and genetic relevance. A user-friendly implementation of this method is available in the survival and multcomp packages of the statistical software R.

Genome-wide association studies involving large population-based samples are used to identify common variants that affect a particular trait. Most of these studies compare the allele frequencies of di-allelic markers in cases and controls using the Cochran-Armitage trend test [1]. Because the mode of inheritance at a given locus is often unknown, a maximum-test (minimum p approach, respectively) based on three mode-specific standardized Cochran Armitage trend tests have been proposed [2]. Alternatively, continuous endpoints (i.e., quantitative traits), such as gene expression, are commonly analyzed using a linear regression model of genotype scores x=(0, 1, 2) adjusted for covariates [3]. A special case of quantitative traits is time-to-event data with right censoring. For example, in a study of the survival of 116 female mice with the three genotypes aa, Aa and AA at the marker DM13D147 in chromosome 13 after an infection with *Listeria monocytogenes* [4], the raw data consist of the following three items (available in the R package qtl [5]): survival time (pheno), genotype group (geno) and censoring status (cens) (Table 1). The related Kaplan-Meier estimators reveal substantial differences in survival between the three genotype groups aa, Aa and AA, in which A is assumed to be the high risk allele:

Figure 1 shows that the survival function of the heterozygous genotype, Aa, is not symmetrical to the functions of the two homozygous genotypes, aa and AA, which would be an indicator of an additive mode of inheritance. Instead, the heterozygous genotype is close to the non-risk homozygous, aa, indicating a recessive mode of inheritance.

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Table 1: Raw data.
The idea of the maximum test is sensitivity to each of the basic modes of inheritance (i.e., additive, recessive, and dominant). Although reporting tiny p-values for the list of top-k SNPs is common nowadays, the ‘Strengthening the Reporting of Genetic Association studies’ report [6] recommended reporting appropriate effect size estimators and their confidence intervals. To compare survival functions, the hazard ratio is the appropriate effect size and simultaneous confidence intervals for a maximum-test for the three basic genetic models (i.e. additive, recessive, and dominant), are estimated. We propose a testing procedure that not only is sensitive to these three alternatives, but also able to determine which of the alternatives is likely using the diagnostic characteristics of simultaneous confidence intervals. We use the multiple contrast test approach [7], extended to the Cox proportional hazard model. This test is not likely applicable to genome-wide studies, multiple contrast test approach [7], extended to the Cox proportional hazard model: (about 0.03 sec on i7-4600 CPU per phenotype and SNP), but it can be used for specific analysis of the top-k SNPs or a priori genes of interest.

Here, we describe an asymptotic multiple contrast test for a censored time-to-event trait assuming the Cox proportional hazards model based on the simultaneous inference approach in general parametric models [8]. This is an extension of a related scores test for the generalized linear model [9] for censored time-to-event traits.

**Methods**

**Marcus-type association test for censored time-to-event data**

We consider three genotype groups \( i \in \{aa, Aa, AA\} \) with \( n_i \) subjects carrying genotype \( i \). A denotes the high risk allele and \( a \) denotes any other allele. To describe the effect of genotype \( i \) and, when applicable, the effects of other covariates on the hazard of death we use a Cox proportional hazard model:

\[
\lambda(t | x_j) = \lambda_0(t) \exp \left( x_j^T \beta \right), j = 1, ..., \sum_{i \in \{aa, Aa, AA\}} n_i.
\]

The vector \( x_j \) contains the covariates of the \( j \)-th individual including the genotype \( i \); the vector \( C = (c_{aa}, c_{Aa}, c_{AA}, 0, ..., 0) \in \mathbb{R}^p \) with restriction \( \sum_{i \in \{aa, Aa, AA\}} \beta_i = 0 \) includes the genotype effects and the effects of further covariates; \( \lambda_0(t) \) denotes the baseline hazard rate at time \( t \) and is assumed to be identical for all individuals.

Let \( C = (c_{aa}, c_{Aa}, c_{AA}, 0, ..., 0) \in \mathbb{R}^p \) be a vector of contrast coefficients fulfilling the constraint \( \sum_{i \in \{aa, Aa, AA\}} c_i = 0 \). For the sake of simplicity, we assume a model with the genotype as single covariate in the following, i.e., \( C=(c_{aa}, c_{Aa}, c_{AA}) \) and \( \beta=(\beta_{aa}, \beta_{Aa}, \beta_{AA}) \). If the elements of vector \( c \) fulfill \( \sum_{i \in \{aa, Aa, AA\}} c_i = -1 \) and \( \sum_{i \in \{aa, Aa, AA\}} c_i = 1 \) the linear combination \( L = \sum_{i \in \{aa, Aa, AA\}} c_i \beta_i \) can be interpreted as a difference of weighted averages of genotype effects.

We consider three genetic contrasts

\[
L_m = \sum_{i \in \{aa, Aa, AA\}} c_m \beta_i, \quad m \in \{\text{dom}, \text{add}, \text{rec}\},
\]

each corresponding to one of the three genetic models, i.e. for each genetic model an individual statistic is computed. These contrasts are formulated by the so-called Marcus-type contrast matrix [10]

\[
C_{\text{Marcus}} = \begin{pmatrix}
  c_{\text{dom}} & c_{\text{add}} & c_{\text{rec}} \\
  n_{aa} & n_{Aa} & n_{AA} \\
  n_{aa} & n_{Aa} & n_{AA} \\
  n_{aa} & n_{Aa} & n_{AA}
\end{pmatrix} \in \mathbb{R}^{3 \times 3},
\]

whose elements \( c_m \) are the contrast coefficients. The product of each row vector of \( C \) and the vector of genotype effects \( \beta=(\beta_{aa}, \beta_{Aa}, \beta_{AA}) \) corresponds to one linear combination \( L_m, \quad m \in \{\text{dom}, \text{add}, \text{rec}\} \) associated with a specific genetic model.

In case of a dominant mode of inheritance, the effects \( \beta_{aa} \) and \( \beta_{Aa} \) on the hazard rate are identical. Therefore a genetic contrast for this mode can be expressed by

\[
L_{\text{dom}} = c_{\text{dom}} \beta = -\beta_{aa} + \frac{n_{aa}}{n_{aa} + N_{aa}} \beta_{aa} + \frac{n_{Aa}}{n_{aa} + N_{aa}} \beta_{Aa} + \frac{n_{AA}}{n_{aa} + N_{aa}} \beta_{AA},
\]

which denotes the difference between the pooled effects of genotypes \( Aa \) and \( AA \). Analogously, in case of a recessive mode of inheritance, the effects \( \beta_{aa} \) and \( \beta_{Aa} \) on the hazard rate are identical. A recessive genetic contrast can be specified by

\[
L_{\text{rec}} = c_{\text{rec}} \beta = -\beta_{aa} + \frac{n_{aa}}{n_{aa} + N_{aa}} \beta_{aa} - \frac{n_{Aa}}{n_{aa} + N_{aa}} \beta_{aa} + \beta_{AA}.
\]

The genetic contrast for an additive model can be expressed by

\[
L_{\text{add}} = c_{\text{add}} \beta = -\beta_{aa} + \beta_{Aa}.
\]

Thus, the case of no global genetic effect is characterized by \( \beta_{aa} = \beta_{Aa} = \beta_{AA} \), or, equivalently \( \bigcap_{m \in \{\text{dom}, \text{add}, \text{rec}\}} L_m < 0 \). We can test for a genetic model by performing a one-sided union-intersection test on the three linear combinations \( L_m \) with control of the FWER over all three contrasts. That is, we test the intersection of the elementary null hypotheses

\[
H_0 : \bigcap_{m \in \{\text{dom}, \text{add}, \text{rec}\}} L_m \leq 0
\]
versus the union of the elementary alternative hypotheses

$$H_1 : \bigcup_{m \in \{\text{dom add, rec}\}} L_m > 0.$$  

This multiple contrast test was already described for normally distributed variables [10]. Therefore, we denote this as the Marcus-type multiple contrast test.

Instead of multiple tests, lower simultaneous confidence intervals for the linear combinations $L_m$ can be used. Exponentiating the confidence limits leads to confidence intervals for $\exp(L_m)$, which can be interpreted as a hazard ratio of the weighted average of the genotype effects. The presence of an association between genotype and trait is indicated if at least one of the three confidence intervals for the hazard ratios $\exp(L_m)$ excludes the value 1.

In other words, the above procedure tests the null hypothesis that no genetic effect exists against the three alternatives that the mode of inheritance is dominant, additive, or recessive.

An adjustment is needed to ensure that the overall hypothesis (no global genetic effect) is tested at level $\alpha$. The three local hypotheses are positively correlated, and this correlation is included in the global genetic effect) is tested at level $\alpha$. The three local hypotheses are

$$\Phi(q) = \mathbb{P}(Z_m \leq q, \forall m \in \{\text{dom add, rec}\}) = 1 - \alpha,$$

where $Z_m$ is the $m$th element of a trivariate normal random vector $Z \sim N(0, R)$. The probability that at least one of the simultaneous confidence intervals does not include the true value of the associated contrast $L_m$ is $\alpha$ with $n \to \infty$. Control of the FWER is achieved using quantiles that take the number of estimated contrasts and correlation between them into account.

Again, exponentiating the lower limit leads to simultaneous confidence intervals for multiple hazard ratios:

$$\exp \left( \sum_{i \in \{\text{aa, Aa, AA}\}} c_i \hat{\beta}_i - z_{3,1-a} \sqrt{\sum_{i \in \{\text{aa, Aa, AA}\}} \sum_{j \in \{\text{aa, Aa, AA}\}} c_i c_j \hat{V}_{ij}} \right),$$

where $z_{3,1-a}$ is the upper equicoordinate $(1-a)$ quantile of the multivariate normal distribution with expectation 0 and correlation matrix $R$ and $\Phi(0, R)$ the associated cumulative density function. The quantile $z_{3,1-a}$ is chosen such that

where $L_m$ is sensitive to three different genetic models and has greater power to

Results

Simulations

To evaluate the performance of the proposed method we estimated the type I error rate and power using simulations in the open-source software R [14]. Each simulation step was repeated 10,000 times.

The trait genotypes for $N=500, 1000, 2000$ subjects were randomly drawn from a multinomial distribution assuming Hardy-Weinberg equilibrium. Allele frequencies were chosen $p=0.5$ at trait locus, $pm=0.05, 0.1, 0.3, 0.5$ at trait marker, and linkage disequilibrium (LD) was chosen $\beta=0.025, 0.05, 0.1, 0.2$. Phenotypic time-to-event data were generated according to Bender et al. [15] using a Weibull distribution with baseline hazard rate $\lambda(t) = \gamma_1 \gamma_2 t - \gamma_1 t^\gamma_2 = 0.01, \gamma_2 = 2$. Censoring times were generated from a uniform distribution on the interval $[0, \tau]$ with the $\tau$ chosen such that the censoring rate was approximately 20%.

The desired confidence level was $1-a=0.95$.

For estimation of the probability of type I error, data were generated under the null hypothesis $\beta_{aa}=\beta_{Aa}=\beta_{AA}=0$, i.e. corresponding to a hazard ratio of 1. In one setting the model was investigated without additional covariates besides the genotype. In another setting, the model was investigated with two covariates: $x$ uniformly distributed on $[2,4]$ without an effect on the hazard rate, i.e. $\hat{\beta}_x=0$, and $x_2$ uniformly distributed on $[0,4]$ with effect $\hat{\beta}_{x_2}=0.5$. The family wise error rate (FWER), that is the probability of falsely detecting any mode of inheritance, was used as measure of the type I error and estimated by the proportion of datasets in which at least one simultaneous confidence interval for Marcus-type hazard ratios did not include the value 1.

For estimation of the power phenotypic values were simulated using genotype effects $\beta_{aa}=\beta_{Aa}=\beta_{AA}=0$ and $\beta_{aa} \in [0,2]$ for a recessive mode of inheritance, $\beta_{aa}=0$, $\beta_{Aa} \in [0,1]$ and $\beta_{aa}=2$ $\beta_{AA}$ for an additive mode of inheritance, and $\beta_{aa}=0$ and $\beta_{AA} \in [0,2]$ for a dominant mode of inheritance. These genotype-specific effects correspond to mode of inheritance-specific hazard ratios $HR \in [1,4.7]$. Each value of power was estimated by the proportion of datasets in which the correct mode of inheritance was detected by the simultaneous lower confidence intervals.
The estimated type I error (FWER) is shown in Figure 2. For fixed sample size, the procedures get more liberal with increasing disbalance of allele frequencies and/or lower LD. For settings with rather balanced allele frequencies, i.e. \(p_m=0.3, 0.5\), a sample size of \(N=500\) is sufficient to ensure FWER control even when LD is low. For settings with unbalanced allele frequencies, i.e. \(p_m=0.05, 0.1\) larger samples are required. A sample size of \(N=2000\) for \(p_m=0.1\), and a sample size of \(N=5000\) for \(p_m=0.05\) provides FWER control (results for \(N=5000\) not shown). In the setting with covariates, the FWER is slightly higher than in the model with the genotype as single covariate.

The power of the procedure to identify the correct genetic model is given in Figure 3 for sample sizes of \(N=1000\). The power increases with higher linkage disequilibrium in all models. The power is considerably higher for the dominant and additive mode of inheritance compared to the recessive mode of inheritance, with the latter showing very poor power except when allele frequency was \(p_m=0.5\) at trait marker. In the dominant model, the power decreases with increasing frequencies of the rare allele, whereas the power increases with increasing frequency of the rare allele in the recessive model. The power in the additive model is similar for all allele frequencies. With increasing sample size (\(N=500\) vs. \(N=1000\) vs. \(N=2000\)), no general increase in the power to detect the correct mode of inheritance can be found (Power curves for \(N=500\) and \(N=2000\) not shown.). In some settings the mode-specific power increases slightly, whereas in some settings the mode-specific power decreases. The power to detect any association increases with increasing sample size, but more often incorrect mode of inheritance is chosen. When the high-risk allele is rare and thus genotype frequencies are unbalanced, an additive mode of inheritance is more often stated to be dominant, and a recessive mode of inheritance is more often stated to be additive (Figure 3).

Figure 4 shows the power of the procedure to detect any association between genotype and trait. The extent to which the procedure’s power to detect the correct mode of inheritance and the power to detect any association differ depends on the underlying mode of inheritance and the disbalance of allele frequencies. A ‘symmetrical’ relation of the modes of inheritance and the differences between overall and mode-specific power exists, which is caused by the fact that the three modes of inheritance are ‘symmetrical’ for the alleles. When high-risk alleles are rare (\(p_m=0.05, 0.1\)), the difference in overall power and mode-specific power is negotiable for the dominant, considerable for the recessive, and intermediate for the additive mode of inheritance. When high-risk alleles are frequent (\(p_m>0.75\)) the difference in overall power and mode-specific power is negotiable for the recessive, considerable for the dominant, and again intermediate for the additive mode of inheritance. When alleles frequencies are balanced, the difference in overall power and mode-specific power is negotiable for the additive model, and intermediate for the dominant and recessive modes of inheritance (Figure 4).

Evaluation of the example

The Listeria example described above was analyzed using the new Marcus-type association test for censored time-to-event data. Simultaneous lower 95% confidence intervals for contrasts of genotype effects corresponding to the three genetic models were computed using the R [14] packages survival [16] and multcomp [17]. The estimated hazard ratios and simultaneous lower confidence limits for the three basic modes of inheritance are given in Table 2.

Clearly, the largest hazard ratio with the most distant lower confidence limit was determined for the recessive mode of inheritance (abbreviated with \(C_{\text{rec}}\)). Mice homozygous for the high risk allele had a
greater chance of a short survival time compared to animals carrying either one or two non-risk alleles (genotypes Aa and aa). With a probability of 95% animals carrying two high risk alleles had at least a 2.23-fold greater hazard.

Software

The simultaneous confidence intervals for hazard ratios obtained from a Cox proportional hazard model can be computed using the coxph function in the package survival [16]. The lower simultaneous confidence intervals for Marcus-type hazard ratios can be computed by the function glht in the package multcomp [17,18]. Intervals for the hazard ratios were obtained by exponentiating the estimated confidence limits. Alternatively, adjusted p-values can be calculated for the corresponding multiple tests. The package multcomp employs the algorithms used for computing the multivariate normal quantiles [19] implemented in the package mvtnorm [20].

Conclusions

Evaluation of selected SNPs by the proposed Marcus-type multiple contrast test for censored time-to-event data in population-based association studies is useful in several aspects. First, the most likely underlying mode of inheritance, not just a global association, can be concluded. The outcomes, i.e., the mode-specific hazard ratios and their lower simultaneous confidence limits, allow interpretation in terms of both statistical significance and medical relevance. This asymptotic approach is limited to study designs with sufficient sample sizes, such as 1000 or more, and is limited to non-rare alleles. Real data can be analyzed easily using the R packages survival and multcomp. Straightforward extensions for data with non-proportional hazards and/or multiple studies (i.e., meta-analysis) are possible.

Acknowledgments

This work was supported in part by HO 16879-1 for the second author (LAH).

References


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Figure 4: Estimated power to detect any association for the three basic genetic models.