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Original Article

Use of multi-state models for time-to-event data

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ABSTRACT

Background & Aim: In medical sciences, the outcome is the time until the occurrence of an event of interest. A multi-state model (MSM) is used to model a process where subjects' transition takes place from one state to the next. For instance, a standard survival curve can be thought of as a simple MSM with two states (alive and dead) and the transition between these two-state models is a method used to analyze time to event data. The most important aspect of this model is that it considers intermediate events and models the effect of covariates on each transition intensity. Some diseases like cancer, human immunodeficiency virus (HIV), etc. have several stages. In the present study, these models were reviewed using cardiac allograft vasculopathy (CAV) data focusing on different approaches.

Methods & Materials: The data of 576 CAVs were collected. A time dependent simple Cox regression model (CRM) was fitted and a three-state illness-death model was considered for the MSMs.

Results: In the simple CRM, only the individuals with the age of > 50 were significant, however for Cox Markov model (CMM) and Cox semi-Markov model (CSMM), the donor's age > 40, sex, and the individuals with the age of > 50 were other significant covariates.

Conclusion: The CMM and CSMM showed more accurate results about risk factors compared to the simple CRM.

Introduction

In medical research, longitudinal datasets have a significant importance. Multi-state models (MSMs) create a feasible framework to analyze multiple records of individuals. Patients experience several stages of a disease over time and every stage has its own features making it important to study with more details. MSMs are useful tools to study the intermediate events in a history of subset of longitudinal data. In situations where estimation of survival probabilities is of interest, it is then a question whether inclusion of information on the

intermediate events in a MSM will give rise to estimators which are in any sense better than what may be obtained from a simple analysis of the marginal distribution of the time to failure (1). Unlike traditional time-to-event analysis where only one outcome is possible for each individual, Markov chains allow analysts to calculate survival times in multiple states. This is particularly attractive for studies of chronic diseases with well-defined phases, like cancer (2) A MSM models a stochastic process with a set of discrete states (at least two). Events are transitions between states, however some states are absorbing, meaning that there is not any transition from them, like death. The simplest including MSM mortality model 2 states and more 1 transition state makes the model complicated. Survival models are capable

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of analyzing incomplete data. Left censor, right censor, interval censor, and left truncation can happen, hence adding more difficulty to analyze the model and construct likelihood function (3). Censoring causes underestimate time to events. In some cases, it was assumed in this study that the censor and truncated data are independent of the process (4). In the present study, a 3-state illness-death (disability) model was considered with 3 transitions, which showed the progression of a disease (Figure 1). In this model, healthy individuals become diseased and then die or healthy individuals die because of other reasons.

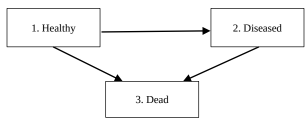


Figure 1. Three state illness-death model

There are two important terms in this context, transition probability and transition intensity or transition rate.

A multi-state process is a stochastic process $(X(t), t \in T)$ with a finite number of states $S = \{1, ..., N\}$.

The time interval is $T=[0,\tau], \tau < \infty$ and state occupied at t is the value of time t. A multi-state process X (.) generates a history $H_{t^-}(\text{6algebra})$ consisting of the observation of the process in the interval $[t,\infty)$. In relative to this history, transition probabilities may be defined by the following relation:

$$\begin{aligned} &p_{hi}(s,t) = P(X(t) = j \mid X(s) = h, Hs -) \\ &\alpha_{hi}(t) = \lim_{\Delta t \to 0} \frac{p_{hi}(t,t+\Delta t)}{\Delta t} \end{aligned}$$

Transition intensity displays potential hazard of progression to state j conditionally on occupying state h.

In this study, the focus is on the Markov and semi-Markov MSMs:

1) *Markov models:* Markov chain models can accommodate censored data, competing risks (informative censoring), multiple outcomes, recurrent outcomes, frailty, and

inconstant survival probabilities (2). Future evolution depends on current state rather than the events occurred before the current state. In most applications, a Markov model is assumed for the MSMs (5). When there are covariates, a Cox (6) model is used to model the effect of covariates on each transition intensity.

2) *Semi-Markov models:* The future evolution of the process does not depend on the current time but rather on the duration in the current state (7).

The goal in the present study was to fit MSMs and time dependent Cox regression model (CRM) on a time-to-event data and compare them.

Methods

The well-known cardiac allograft vasculopathy (CAV) dataset studied by Sharples et al. (8) were used in the present study. This dataset participants who received included 576 transplants between august 1979 to January 2000 and survived at least 1 year after transplantation and had undergone at least one coronary angiogram. The state at each time was a grade of CAV, a deterioration of the arterial walls (9). Approximately, each year after transplantation, each patient had an angiogram, at which CAV could be diagnosed. The result of the test was in the variable state, with possible values 1 and 2, representing CAV-free and CAV, respectively. A value of 3 was recorded at the date of death. Years designated the time of the test in years since the heart transplantation. Other variables included age (age at screen), donor age, and sex (0 and 1 representing men and women, respectively). Age and donor age were considered categorical, however the sex variable was dichotomous. Two cut points for age and donor's age were taken into account mostly based on the quantiles. 30 and 50 were the cut points of age variable and for 20 and 40.

Time dependent CRM: In the analysis of time-to-event data, the Cox proportional-hazards regression model has achieved popularity due to considering the censoring and the covariates. The Cox model with time-dependent covariate was much more complex than with fixed (time

invariant) covariates due to the possibility of changing the values over time. This model involved constructing a function of time (10). The marginal distribution of survival times, that is, without consideration of CAV, was analyzed using the Cox proportional hazards regression model (6). Thus, the hazard function for individual i with covariates Z_i was assumed to have the following form with unspecified baseline hazard function $\alpha_0(t)$ and regression coefficients β .

$$\alpha_i(t; Z_i) = \alpha_0(t) \exp(\beta_i' Z)$$

The survival probability $\hat{S}(t; Z)$ for given covariates of simple Cox model, Z, maybe estimated by

$$\widehat{S}(t; Z) = \prod_{u \le t} (1 - d\widehat{A}_0(u) \exp(\widehat{\beta}Z))$$

Where, $\widehat{A}_0(t)$ is the Breslow estimator for the integrated baseline hazard function.

Cox-Markov model (CMM): Finding a relation between predictor variables and the response variable was one of the biggest goals of MSMs in the present study. This model have been used in literature to relate the individual characteristics to the intensity transition through a possibly time-dependent covariate vector Z (11). The simplest way was to decouple the whole process into several survival models. The illness-death model has been presented in figure 2, fitting separate intensity rates to all possible transitions using semi-parametric proportional hazard regression models, while making appropriate adjustments to the risk set (7). Using Cox-like models of the form $\alpha_{hi}(t; Z) = \alpha_{hi,0}(t) \exp(\beta_{hi}^T Z),$ the transition intensities for illness death model $\alpha_{hi}(t; \mathbb{Z}), 1 \le h < j \le 3$, (4). This model was CMM which assumes the process to be Markovian. Estimating transition probability, taking into account independent z variables, yields:

$$\begin{split} & p_{hj}(\,s,t\mid Z\,) = P(X(t) = j\mid X(s) = h,Z) \\ & h \leq j \qquad S \leq t \\ & \hat{p}_{11}(\,s,t\mid Z\,) = \prod_{s < u \leq t} (1 - \sum_{j=2}^3 d\widehat{A}_{1j}(u\mid Z) \\ & \hat{p}_{22}(\,s,t\mid Z\,) = \prod_{s < u \leq t} (1 - d\widehat{A}_{23}(u\mid Z) \end{split}$$

$$\hat{p}_{12}(s, t \mid Z) = \prod_{u < t} \hat{p}_{11}(s - u \mid Z) d\hat{A}_{12}(u \mid Z) \hat{p}_{22}(u +, t \mid Z)$$

Cumulative hazard function was $\widehat{A}_{hj}(t \mid Z) = \widehat{A}_{hj0}(t) \exp(\widehat{\beta}_{hj}Z)$ using $\widehat{A}_{hj0}(t) = \int_0^t \alpha_{hj0}(u)$ to estimate $\widehat{A}_{hj0}(.)$.

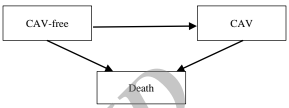


Figure 2. A three-state model for the cardiac allograft vasculopathy (CAV) data

Cox semi-Markov model (CSMM): Markov models have some limitations which hindered its use. Markov assumption was violated in some situations, meaning that the future depended on the individual's past only through his/her current state. In this model, the time was reset to zero when a new state was reached, hence the time scale was called "clock reset". Each time the patient entered a new state, time was reset to 0. Therefore, CSMM could be easily fitted for the illness-death model; transition $2 \rightarrow 3$ was the difference between CMM and CSMM. The Markov assumption was not satisfied, as the mortality after getting CAV depended strongly on the time since getting CAV, suggesting that a semi-Markov model might be more appropriate (12). It was only needed to remodel α_{23} , so that it depended only on the duration in state 2. Using time since getting CAV as the basic time scale for the model for mortality after getting CAV, We model α_{23} as follows

$$\begin{aligned} \alpha_{23}(t-t_{2i};Z) &= \\ \alpha_{230}(t-t_{2i}) \exp\left(\beta_0^T Z_{23i}\right) & t \geq t_{2i} \end{aligned}$$

Where, t_{2i} was the entry time into state 2. R package p3state.msm was used for analyses. The significance level was considered to be 0.050 in this study.

Results

15% and 85% of the participants were women and men, respectively. The mean age of the

individuals was 47 (47 and 43 for men and women, respectively). The mean age of donors was 30. The number of individuals suffering from CAV was 179 out of 576 (31.00%). Among the patients, 12 (14.46%) women and 167 (33.87%) men had CAV. Further information was listed in tables 1 and 2. During the study period, 576 individuals were at risk of transition from the state 1 to states 2 and 3 and transition from the state 2 to state 3. Of these, 179 (167 and 12 men and women, respectively) experienced a transition to the state 2, and 139 (120 and 19 men and women, respectively) to state 3.

Table 1. Continuous characteristics of the study variables

	Valid	Age	Donor's age
N	vanu	576	576
	Missing	0	0
Mean		47.21	30.78
Mean SE		0.45	0.51
Median		49.61	29.00
Mode		41.80	18.00
SD		10.85	12.29
Variance		117.77	151.11
Range		57.96	61.00
Minimum		6.30	0.00
Maximum		64.26	61.00
Sum		27192.28	17727.00
Percentiles	25	42.29	20.00
	50	49.61	29.00
	75	55.21	40.75

SE: Standard error; SD: Standard deviation

95 individuals (88 and 7 men and women, respectively) died without exposing to CAV. The number of individuals remaining in state 1 was 258. In addition, 84 of individuals were censored on transition from state 2.

The transition probabilities are changing over time. In the present study, transition probabilities were calculated only for median of time in table 3, the probability of remaining CAV-free cases was 0.78. The estimated probability for individuals with transition from state 1 to state 2 was 0.14. For patients with CAV, there was a probability of 0.18 to die and the probability of dying because of reasons other than CAV was 0.08. Patients with CAV had a 0.82 probability to remain in state 2. It is noteworthy that these estimated transition probabilities were only valuable at median of time and they will change for other values of time (13).

Model fit

Time dependent CRM: In case of ignorance of the intermediate event, the results with time dependent CRM shown in table 4 indicated that only the individuals > 50 years old are significant (P = 0.0027). The individuals > 50 years old have 2.46 chance of death compared with those with < 30 years old.

Multi-State Models (MSMs)

Cox Markov Model (CMM): In transition from the state 1 to the state 2 as see in table 5, donor's age variable for 20-40 years old donors was in borderline significance (P = 0.053), and for more than 40 years old, donors were highly significant (P < 0.001). Individuals with donors > 40 years old had a 2.233 chance of CAV compared to individuals with donors < 20 year old. The sex variable was significant as well (P = 0.030). Comparing to men, women had 0.518 hazard of CAV.

In transition from the state 1 to the state 3 in table 6, it was shown that the age variable for category of > 50 years old CAV-free individuals was significant (P < 0.001). Individuals > 50 years old had 3.652 chance of dying without getting CAV compared to < 30 year-old individuals.

Table 2. Table of frequencies

		Without CAV		CA	V
		Alive rate (%)	Dead rate (%)	Alive rate (%)	Dead rate (%)
Car	Men	206 (63.2)	120 (36.8)	79 (47.3)	88 (52.7)
Sex Women	52 (73.2)	19 (26.8)	5 (41.7)	7 (58.3)	
	< 30	27 (73.0)	10 (27.0)	8 (36.4)	14 (63.6)
Age (year)	30-50	99 (67.8)	47 (32.2)	41 (41.8)	57 (58.2)
	> 50	132 (61.7)	82 (38.3)	35 (59.3)	24 (40.7)
Donorlo ano	< 20	58 (65.2)	31 (34.8)	13 (36.1)	23 (63.9)
Donor's age	20-40	136 (66.7)	68 (33.3)	39 (41.1)	56 (58.9)
(year)	> 40	64 (61.5)	40 (38.5)	32 (66.7)	16 (33.3)

CAV: Cardiac allograft vasculopathy

Table 3. Transition probabilities for median of time

	State 1	State 2	State 3
State 1	0.78	0.14	0.08
State 2	0	0.82	0.18

The donor's age > 40 years old was an important risk factor (P = 0.043). In addition, individuals with donor of > 40 years old had 1.665 chance of dying without getting CAV comparing to individuals with donor age of < 20 years old. Moreover, in transition from the state 2 to the state 3, there was no important risk factor depending on CMM.

Table 4. Time-dependent Cox regression model (CRM)

	Coef	Exp (coef)	Se (coef)	z	P-value	Lower	Upper
Factor (age) 1	0.131	1.140	0.238	0.549	0.583	0.714	1.818
Factor (age) 2	0.901	2.463	0.247	3.646	0.000	1.517	3.998
Factor (dage) 1	0.076	1.079	0.170	0.445	0.657	0.773	1.505
Factor (dage) 2	0.201	1.222	0.201	1.000	0.317	0.825	1.811
Sex	0.399	1.490	0.218	1.831	0.067	0.972	2.283
Treat	1.013	2.754	0.160	6.340	< 0.001	2.014	3.767

Likelihood ratio test = 69.01099 on 6 df, P < 0.001

Cox Semi Markov Model (CSMM): Markov assumption should only check for transition from 2 to 3 in the illness-death model. The Markov assumption was highly significant in the present study (P = 2.27e-11).

Table 5. Cox Markov Model (CMM) transition from state 1 to state 2

	Coef	Exp (coef)	Se (coef)	Z	P-value	1	u
Factor (age) 1	0.189	1.208	0.242	0.780	0.435	0.751	1.943
Factor (age) 2	-0.142	0.867	0.261	-0.546	0.585	0.520	1.446
Factor (dage) 1	0.390	1.476	0.202	1.928	0.054	0.993	2.194
Factor (dage) 2	0.803	2.233	0.229	3.515	< 0.001	1.427	3.495
Sex	-0.657	0.519	0.304	-2.161	0.031	0.286	0.941

dage: Donor's age

Likelihood ratio test = 22.85106 on 5 df, P < 0.001

Table 6. Cox Markov Model (CMM) transition from state 2 to state 3

	Coef	Exp (coef)	Se (coef)	z	P-value	l	u
Factor (age) 1	0.436	1.547	0.359	1.216	0.224	0.766	3.127
Factor (age) 2	1.295	3.652	0.357	3.632	< 0.001	1.815	7.347
Factor (dage) 1	0.164	1.178	0.224	0.732	0.464	0.759	1.829
Factor (dage) 2	0.510	1.665	0.252	2.025	0.043	1.017	2.726
Sex	0.368	1.445	0.256	1.436	0.151	0.874	2.389

dage: Donor's age

Likelihood ratio test = 37.28319 on 5 df, P < 0.001

This assumption was tested in table 7, showing that the transition rates in states were affected by the previous sojourn time. Therefore, CSMMs could be a better choice. This model was fitted only for transition from the state 2 to the state 3. Results in table 8 indicated that the donor's age > 40 years old was significant (P = 0.034). Patients with CAV with donor's age of > 40 year old had 0.48 chance of death compared to the patients with CAV with donor's of < 20 years old.

 Table 7. Markov assumption test

	Coef	Exp (coef)	Se (coef)	Z	P-value
Factor	-0.398	0.672	0.330	-1.210	0.227
(age) 1 Factor (age) 2	0.280	1.323	0.359	0.780	0.435
Factor (dage) 1	-0.238	0.788	0.284	-0.840	0.400
Factor (dage) 2	-0.753	0.471	0.356	-2.110	0.035
Sex Start	0.787 -0.292	2.197 0.747	0.409 0.046	1.920 -6.280	0.054 < 0.001

dage: Donor's age

Likelihood ratio test = 61.5 on 6 df, P < 0.001

n = 179, number of events = 95

Discussion

In the present study, the data were analyzed concerning CAV and mortality among patients with transplantation with three-state illness-death models obtained by splitting the alive state from the simple two-state model for survival data into the states of healthy heart transplant recipients, CAV, and death.

^{-2 *} Log-likelihood = 2358.093

dage: Donor's age

^{-2 *} Log-likelihood = 1904.55

^{-2 *} Log-likelihood= 1444.886

Table 8. Cox Semi-Markov Model (CSMM) from state 2 to state 3

	Coef	Exp (coef)	Se (coef)	Z	P-value	l	u
Factor (age) 1	-0.313	0.731	0.331	-0.945	0.345	0.382	1.400
Factor (age) 2	0.196	1.216	0.364	0.538	0.590	0.596	2.483
Factor (dage) 1	-0.291	0.748	0.276	-1.055	0.292	0.436	1.284
Factor (dage) 2	-0.734	0.480	0.347	-2.116	0.034	0.243	0.947
Sex	0.653	1.921	0.400	1.631	0.103	0.877	4.211

dage: Donor's age

Likelihood ratio test = 9.381181 on 5 df, P = 0.095

In this study, the use of MSMs in the analysis of survival data were discussed in the scope of the three-state model. Nonparametric estimators for the transition probabilities have been presented and illustrated using a real dataset on CAV. The first model, a Markov process model did not fit the data since the mortality after CAV depended strongly on the time since CAV, and, instead, semi-Markov model was studied. These three-state models turned out to provide important clinical information by highlighting covariates affecting the mortality and covariates affecting the CAV. The main message of this study, however, is the fact that the precision of survival probability estimates based on the three-state models tended to be better than for those based on the simple Cox model. As it can be seen in the results, there are different risk factors affecting every single transitions. However, the result of simple CRM is not precise as MSMs.

Conclusion

In conclusion, the present study has shown that a MSM may in some cases be preferable to a model for the marginal survival distribution. Thus, for the MSM to be analyzed simply, exact times of transition are required when the observation of some transition times are interval-censored, hence the likelihood expressions become more involved.

Moreover, the multi-state analysis requires some assumptions concerning a Markov or semi-Markov structure of the data assumptions which are avoided in the marginal analysis (14). Final problem with the MSMs is that estimates for transition probabilities are not generally available for semi-Markov models. At last, it is important to realize that there are several

modelling strategies for the intermediate event and that each of these strategies has restrictions and may cause different results.

Conflict of Interests

Authors have no conflict of interests.

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^{-2 *} Log-likelihood = 708.6002

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