# Modeling the cumulative incidence function of clustered competing risk data





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# Clustered competing risk data

#### Key terms:

- » Clustered: groups with a dependence structure (e.g. families);
- » Causes competing by something.

#### Something?

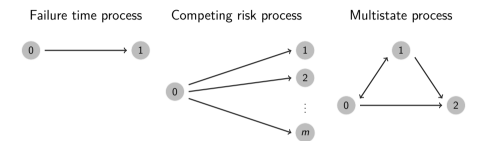
- » Failure of an industrial or electronic component;
- » Occurence or cure of a disease or some biological process;
- » **Progress** of a patient clinic state.

Independent of the application, always the same framework

Group	ID	Cause 1	Cause 2	Censorship	Time	Feature
1	1	Yes	No	No	10	Α
1	2	No	No	Yes	8	Α
2	1	No	No	Yes	7	В
2	2	No	Yes	No	5	Α

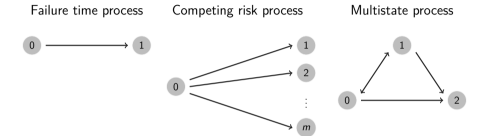


# Data designs





## Data designs



#### Modeling framework

We have to choose which scale we model the survival experience.

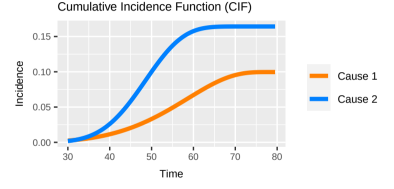
Usually, is the

hazard (failure rate) scale :  $\lambda(t \mid \text{features}) = \lambda_0(t) \times c(\text{features})$ .



#### In the competing risk setting ...

a more attractive possibility is to work on the probability scale, focusing on the cause-specific

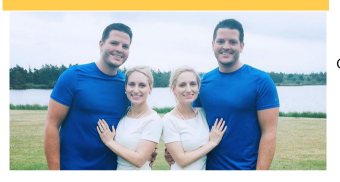


i.e.

 $\mathsf{CIF} = \mathbb{P}[\mathsf{\ failure\ time} \leq t, \mathsf{\ a\ given\ cause} \mid \mathsf{features\ }]$ 



## Main focus application: cancer incidence in twins



Clustered competing risks data

Lack Clusters? Families

Lack Family studies

Lack Twins data

Family studies  $\Rightarrow$  within-family dependence

#### That may reflect

- » Disease heritability;
- » The impact of shared environmental effects;
  - » Parental effects: continuity of the phenotype across generations.



# Our contribution: a hierarchical approach

#### Thinking on two competing causes

... for the outcome  $y_{ijt}$  of a subject i, family j, in the time t, we have

$$y_{ijt} \mid \underbrace{\left\{u_{1j}, u_{2j}, \eta_{1j}, \eta_{2j}\right\}}_{\text{latent effects}} \sim \text{Multinomial}(p_{1ijt}, p_{2ijt}, p_{3ijt})$$

$$\begin{bmatrix} u_{1j} \\ u_{2j} \\ \eta_{1j} \\ \eta_{2j} \end{bmatrix} \sim \begin{array}{l} \text{Multivariate} \\ \text{Normal} \end{array} \begin{pmatrix} \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \end{bmatrix}, \begin{bmatrix} \sigma_{u_1}^2 & \sigma_{u_1, u_2} & \sigma_{u_1, \eta_1} & \sigma_{u_1, \eta_2} \\ \sigma_{u_2}^2 & \sigma_{u_2, \eta_1} & \sigma_{u_2, \eta_2} \\ \sigma_{\eta_1}^2 & \sigma_{\eta_1, \eta_2} \\ \sigma_{\eta_2}^2 \end{bmatrix} \end{pmatrix}$$

$$p_{kijt} = \frac{\partial \text{CIF}}{\partial t}$$

$$= \frac{\partial}{\partial t} \underbrace{\pi_k(X, u_1, u_2 \mid \beta)}_{\text{pluster accificent of the problem}} \Phi[w_k g(t) - X^\top \gamma_k - \eta_k],$$

risk level

cluster-specific

cluster-specific

failure time trajectory



#### Contributions & challenges

- » A clear and simpler modeling structure;
- » There is no free lunch Computational challenges overcame via an efficient implementation and estimation routines;
- The data is very simple, we just know the outcome (yes or no);
- We have to be able to build the CIF curves;
- » And accommodate the within-family dependence properly, that can happen in different manners;
- **>>** . . .



## Thank you







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