Homework 2

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Due date: Thursday, October 11

1. Show that (algebraically) in the absence of censoring $\hat{S}_{\text{KM}}(t) = \hat{S}_{\text{e}}(t)$. Assume $m = \max\{i : t_{(i)} \leq t\}$, then:

$$\begin{split} \hat{S}_{\text{KM}}(t) &= \prod_{\substack{t_{(i)} \leq t \\ n_i}} \frac{n_i - d_i}{n_i} = \frac{n - d_1}{n} \cdot \frac{n - d_1 - d_2}{n - d_1} \cdot \dots \cdot \frac{n - d_1 - d_2 - \dots - d_m}{n - d_1 - d_2 - \dots - d_{m-1}} = \frac{n - d_1 - d_2 - \dots - d_m}{n} \\ &= \frac{\# individuals \ with \ survival \ times \geq t}{\# individuals \ in \ the \ data \ set} = \hat{S}_{\text{e}}(t) \end{split}$$

2. In the absence of censoring, show that the Greenwood Formula (page 30 on note 2) can be reduced to

$$\frac{\hat{S}_{\text{\tiny KM}}(t) \times \{1 - \hat{S}_{\text{\tiny KM}}(t)\}}{n}.$$

You might assume there are no ties among the observations.

$$\begin{split} &Var\{\widehat{S}_{KM}(t)\}\approx\widehat{S}_{KM}^{2}(t)\cdot\sum_{t_{(i)}\leq t}\frac{d_{i}}{n_{i}\cdot(n_{i}-d_{i})}\\ &=(\frac{n-d_{1}-d_{2}-\cdots-d_{m}}{n})^{2}\cdot\sum_{t_{(i)}\leq t}\frac{d_{i}}{n_{i}\cdot(n_{i}-d_{i})}\quad assume\ no\ censoring,\ follow\ from\ Q1\\ &=(\frac{n-t_{(m)}}{n})^{2}\cdot\sum_{t_{(i)}\leq t}\frac{1}{n_{i}\cdot(n_{i}-1)}\quad assume\ no\ ties\\ &=(\frac{n-t_{(m)}}{n})^{2}\cdot(\frac{1}{n(n-1)}+\frac{1}{(n-1)(n-2)}+\cdots+\frac{1}{(n-(t_{(m)}-1))(n-t_{(m)})})\quad assume\ no\ censoring\\ &=(\frac{n-t_{(m)}}{n})^{2}\cdot((-\frac{1}{n}+\frac{1}{n-1})+(-\frac{1}{n-1}+\frac{1}{n-2})+\cdots+(-\frac{1}{n-(t_{(m)}-1)}+\frac{1}{n-t_{(m)}}))\\ &=(\frac{n-t_{(m)}}{n})^{2}\cdot(\frac{1}{n-t_{(m)}}-\frac{1}{n})\\ &=\frac{n-t_{(m)}}{n}-(\frac{n-t_{(m)}}{n})^{2}}{n}\\ &=\frac{\widehat{S}_{KM}(t)\cdot\widehat{S}_{KM}(t)}{n} \end{split}$$

- 3. Consider the Leukemia data from the survival package:
 - > library(survival)
 - > head(aml)

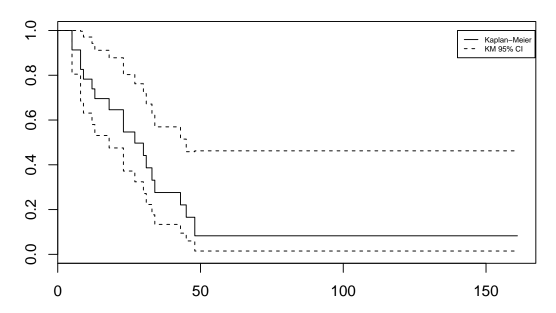
```
time status x
1 9 1 Maintained
2 13 1 Maintained
3 13 0 Maintained
4 18 1 Maintained
5 23 1 Maintained
6 28 0 Maintained
```

In here, each row represent one patient. aml is the observed survival time, status is the censoring indicator (1 = event, 0 = censored), and x is the treatment indicator. We will ignore the treatment indicator for now.

a. Plot the Kaplan-Meier survival curve for the data.

```
> km <- survfit(Surv(time, status) ~ 1, data = aml)
> plot(km,main="Kaplan-Meier survival curve")
> legend(140, 1, legend=c("Kaplan-Meier","KM 95% CI"),col=c("black","black"),
+ lty=c(1,2), cex=0.5)
```

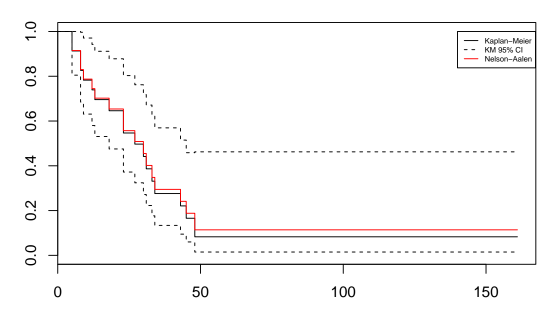
Kaplan-Meier survival curve



b. Add the Nelson-Aalen survival curve to the Kaplan-Meier plot from (3a).

```
> cox <- coxph(Surv(time, status) ~ 1, data = aml)
> H0 <- basehaz(cox)
> plot(km,main="Kaplan-Meier and Nelson-Aalen survival curve")
> lines(H0$time, exp(-H0$hazard), 's', col = 2)
> legend(140, 1, legend=c("Kaplan-Meier","KM 95% CI", "Nelson-Aalen"),
+ col=c("black","black", "red"),lty=c(1,2,1), cex=0.5)
```

Kaplan-Meier and Nelson-Aalen survival curve



4. The expected survival time for the Leukemia data in #(3) does not exist because the last observation is a censored event. An alternative is to lookInstead of looking at the expected survival time, an alternative is to look at the restricted mean survival time. Compute E(T|T<161) based on the survival curve in (3a).

```
> matrix(c(km$time,km$surv),nrow=length(km$time),
+ dimnames = list(NULL, c("death_time","km_surv"))) %>%
+ as.data.frame() %>%
+ add_row(death_time = 0, km_surv = 1,.before = 1) %>%
+ arrange(desc(km_surv)) %>%
+ mutate(time_diff=lead(death_time,default = 0)-death_time) %>%
+ filter(death_time<161) %>%
+ summarise(expected_lifetime=sum(time_diff*km_surv))
```

```
expected_lifetime
1 36.36439
```

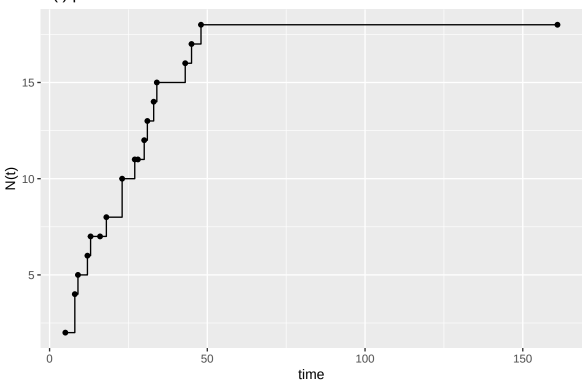
The expected lifetime is 36.36439.

- 5. Let $N_i(t)$ be the number of events over time interval (0,t] for the *i*th patient in #(3). Let $N(t) = \sum_{i=1}^{n} N_i(t)$ be the aggregated counting process.
 - a. Plot N(t).

```
> calculate_Nt=
+ aml %>% arrange(time) %>% select(-x) %>% group_by(time) %>%
+ summarize(di = sum(status)) %>%
```

```
+ mutate(Nt = cumsum(di))
>
> qplot(time, Nt, data = calculate_Nt, geom = "step",
+ main="N(t) plot") +
+ ylab("N(t)") + geom_point()
```

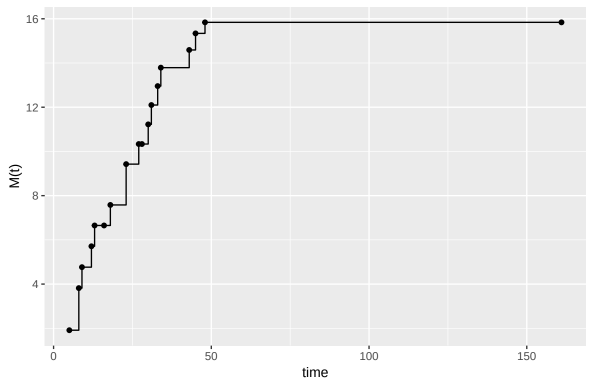
N(t) plot



- b. Plot M(t), where $M(t) = N(t) \hat{H}(t)$ and $\hat{H}(t)$ is the Nelson-Aalen estimator for the cumulative hazard function.
- Note on 5b: After giving some thought, I think it is more meaningful to plot $dM(t) = dN(t) \hat{h}(t)dt$. Both plots will receive full credit for 5b.

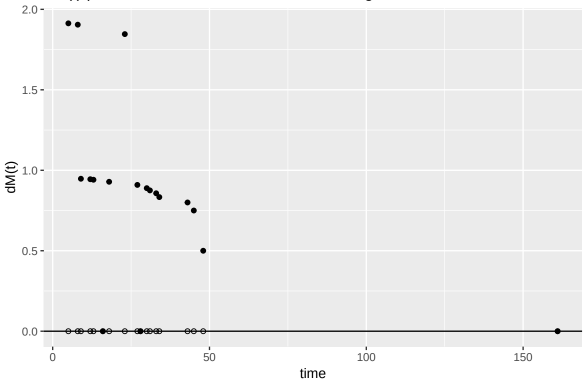
```
> na_surv_by_formula=
+ aml %>% arrange(time) %>% select(-x) %>% group_by(time) %>%
+ summarize(di = sum(status), ni = length(status)) %>%
+ mutate(ni = rev(cumsum(rev(ni))), hi = di/ni,
+ Nt = cumsum(di), Ht = cumsum(hi), Mt = Nt-Ht,
+ dNt = di, htdt = hi, dMt = di-hi)
> qplot(time, Mt, data = na_surv_by_formula, geom = "step",
+ main="M(t) plot from Nelson-Aalen estimator using the formula") +
+ ylab("M(t)") + geom_point()
```

M(t) plot from Nelson-Aalen estimator using the formula



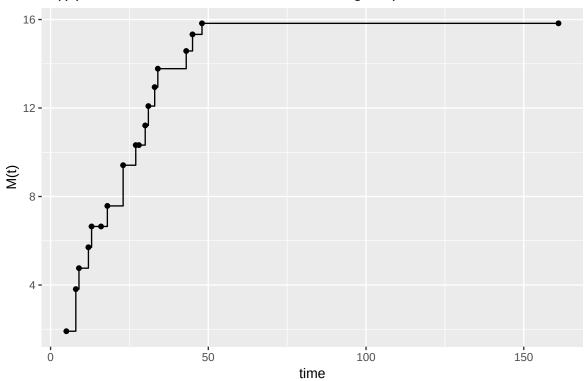
```
> qplot(time, dMt, data = na_surv_by_formula,
+ main="dM(t) plot from Nelson-Aalen estimator using the formula") +
+ geom_abline(intercept = 0, slope = 0) +
+ geom_point(aes(na_surv_by_formula$time, rep(0,18)), pch=1) + ylab("dM(t)")
```

dM(t) plot from Nelson-Aalen estimator using the formula

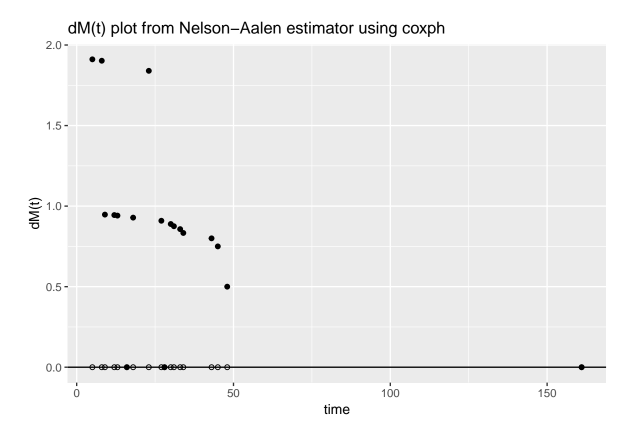


```
> na_surv_by_cosph=
+ aml %>% arrange(time) %>% select(-x) %>% group_by(time) %>%
+ summarize(di = sum(status), ni = length(status)) %>%
+ mutate(ni = rev(cumsum(rev(ni))), hi = di/ni,
+ Nt = cumsum(di), Ht = HO$hazard, Mt = Nt-Ht,
+ dNt = di, htdt = diff(c(0,Ht)), dMt = di-htdt)
>
> qplot(time, Mt, data = na_surv_by_cosph, geom = "step",
+ main="M(t) plot from Nelson-Aalen estimator using coxph") +
+ ylab("M(t)") + geom_point()
```

M(t) plot from Nelson-Aalen estimator using coxph



```
> qplot(time, dMt, data = na_surv_by_cosph,
+ main="dM(t) plot from Nelson-Aalen estimator using coxph") +
+ geom_abline(intercept = 0, slope = 0) +
+ geom_point(aes(na_surv_by_cosph$time, rep(0,18)), pch=1) + ylab("dM(t)")
```



Since the estimators from the formula and the coxph are very close, they plots are almost identical.