# Supplement to: Multiple imputation of missing covariates when using the Fine–Gray model

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## S1 Minimal code example

This is the minimal R code companion to section 3.4 of main manuscript. The parameters from the simulation study scenario with p = 0.15, random censoring, and correctly specified Fine–Gray were used to generate the example dataset below.

```
# Load libraries
library(data.table)
library(survival)
library(kmi)
library(mice)
library(smcfcs)

# Minimal dataset
head(dat, n = 10)
```

```
time D
                     Χ
                             Ζ
   id
1
    1 0.491195 0
                     1
                        0.126
2
    2 0.028680 2 <NA>
                        1.266
3
    3 0.910797 0
                     0 - 1.571
4
    4 0.217566 2
                     1 - 0.500
5
    5 0.132420 2
                     0 0.781
    6 0.800913 2
                     0 - 0.434
6
7
    7 0.041653 2 <NA> -0.844
    8 0.036202 1 <NA>
```

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```
9 9 0.046798 0 0 -1.653

10 10 0.997413 0 <NA> -1.196

sapply(dat, class)

id time D X Z

"integer" "numeric" "factor" "factor" "numeric"

nrow(dat)
```

#### [1] 2000

1. Add columns  $\hat{H}_1(T)$  and  $\hat{H}_2(T)$  to the original data, which are the marginal cause-specific cumulative hazards for each competing risk evaluated at an individual's event or censoring time (obtained using the Nelson–Aalen estimator).

```
# Add cause-specific event indicators + cumulative hazards
dat$D1 <- as.numeric(dat$D == 1)
dat$D2 <- as.numeric(dat$D == 2)
dat$H1 <- nelsonaalen(data = dat, timevar = "time", statusvar = "D1")
dat$H2 <- nelsonaalen(data = dat, timevar = "time", statusvar = "D2")</pre>
```

2. Multiply impute the potential censoring for those failing from cause 2 using  $\{kmi\}$ , yielding m censoring complete datasets (i.e. with "complete" V). Any completely observed covariates that are known to affect the probability of being censored should be included as predictors in the model for the censoring process.  $\{kmi\}$  imputes based on stratified Kaplan–Meier when Z are categorical, and based on a Cox model when at least one of Z are continuous.

```
# 5 imputed datasets
M <- 5

# Multiply impute the censoring times
cens_imps <- kmi(
   formula = Surv(time, D != 0) ~ 1, # Additional predictors added here
   data = dat,
   etype = D,
   failcode = 1, # Specify event of interest
   nimp = M
)</pre>
```

3. In each censoring complete dataset, add an additional column  $\Lambda_1(V)$ . This takes the value of the marginal cumulative subdistribution hazard for cause 1 at an individual's observed or imputed subdistribution time, obtained with the Nelson–Aalen estimator based on I(D=1) and imputed V.

```
# Preparation for covariate imputation:
# Create list of censoring complete datasets (with imputed V)
list_to_impute <- lapply(cens_imps$imputed.data, function(imp_dat) {
    # Adjust new ordering from kmi (cause 2 individuals appended at bottom)
    dat_to_impute <- cbind(cens_imps$original.data, imp_dat)</pre>
```

```
# Compute/add Lambda 1(V) in each imputed dataset
 dat to impute$Lambda1 <- nelsonaalen(</pre>
   data = dat to impute,
   timevar = "newtimes", # kmi naming for V
   statusvar = "D1" # I(D=1)
 return(dat to impute)
})
# newevent is equal to I(D=1)
head(list to impute[[1]])
   id
         time D
                         Z D1 D2
                                         H1
                                                   H2 newtimes newevent
1
   1 0.491195 0
                    3 0.910797 0
                   0 -1.571 0 0 0.25761243 0.83833716 0.910797
                                                                      0
   8 0.036202 1 <NA> 1.564 1 0 0.02028935 0.09603222 0.036202
                                                                      1
   9 0.046798 0
                   0 -1.653 0 0 0.02606228 0.10990397 0.046798
                                                                      0
10 10 0.997413 0 <NA> -1.196 0 0 0.27549886 0.87116320 0.997413
                                                                      0
12 12 0.056015 0 <NA> 0.058 0 0 0.02903112 0.12350351 0.056015
                                                                      0
     Lambda1
1 0.12385222
3 0.16659793
8 0.01932257
9 0.02452308
10 0.17340532
12 0.02715245
```

4. In each censoring complete dataset (each with different V and  $\hat{\Lambda}_1(V)$ , but same  $\hat{H}_1(T)$  and  $\hat{H}_2(T)$ ), create a single imputed dataset using the desired covariate imputation method(s).

```
# Prepare predictor matrices for MICE using first censoring complete dataset
predmat_cs_approx <- predmat_fg_approx <- mice::make.predictorMatrix(
    data = list_to_impute[[1]]
)
predmat_cs_approx[] <- predmat_fg_approx[] <- 0

# Explicitly specify predictors to include in the imputation model
predmat_cs_approx["X", c("Z", "D1", "D2", "H1", "H2")] <- 1
predmat_fg_approx["X", c("Z", "D1", "Lambda1")] <- 1
predmat_fg_approx</pre>
```

	id	time	D	Χ	Z	D1	D2	H1	Н2	newtimes	${\tt newevent}$	Lambda1
id	0	0	0	0	0	0	0	0	0	0	0	0
time	0	0	0	0	0	0	0	0	0	0	0	0
D	0	0	0	0	0	0	0	0	0	0	0	0
X	0	0	0	0	1	1	0	0	0	0	0	1
Z	0	0	0	0	0	0	0	0	0	0	0	0
D1	0	0	0	0	0	0	0	0	0	0	0	0

```
D2
          0
               0 0 0 0 0 0 0
                                          0
                                                   0
                                                            0
          0
               00000000
H1
                                          0
                                                   0
                                                            0
H2
          0
               00000000
                                          0
                                                   0
                                                            0
newtimes 0
               00000000
                                          0
                                                   0
               00000000
                                          0
                                                   0
                                                            0
newevent
          0
               00000000
Lambda1
                                          0
# Prepare the methods:
# - Approx methods: model type for X | Z, outcome
methods_approx <- mice::make.method(data = list_to_impute[[1]])</pre>
# - SMC methods: proposal model for X | Z (need to use {smcfcs} naming)
methods_smcfcs <- mice::make.method(</pre>
  data = list to impute[[1]],
  defaultMethod = c("norm", "logreg", "mlogit", "podds")
methods smcfcs
      id
             time
                                  Χ
                                           Ζ
                                                   D1
                                                             D2
                                                                      H1
      11 11
                        "" "logreg"
                                           11 11
                                                    11 11
                                                             11 11
                                                                      11 11
      H2 newtimes newevent Lambda1
               11 11
# Impute X in each censoring complete dataset
# (parallelise this loop for speed improvements on larger data)
list_imps <- lapply(list_to_impute, function(imp_dat) {</pre>
 m < -1
  iters <- 10
  imps_cs_approx <- mice(</pre>
   data = imp_dat,
   m = m,
   maxit = iters,
   method = methods_approx,
   predictorMatrix = predmat cs approx
  imps_fg_approx <- mice(</pre>
   data = imp dat,
   m = m,
   maxit = iters,
   method = methods approx,
   predictorMatrix = predmat_fg_approx
  imps cs smc <- smcfcs(</pre>
   originaldata = imp_dat,
   smtype = "compet",
```

```
smformula = list(
      "Surv(time, D == 1) \sim X + Z",
      "Surv(time, D == 2) ~ X + Z"
    ),
    method = methods_smcfcs,
    m = m,
    numit = iters
  )
  imps_fg_smc <- smcfcs(</pre>
    originaldata = imp dat,
    smtype = "coxph",
    smformula = "Surv(newtimes, D1) ~ X + Z",
    method = methods_smcfcs,
    m = m,
    numit = iters
  )
  # Bring all the imputed datasets together
  imps <- rbind.data.frame(</pre>
    cbind(method = "CCA", imp_dat),
    cbind(method = "cs smc", imps cs smc$impDatasets[[1]]),
    cbind(method = "cs_approx", complete(imps_cs_approx, action = 1L)),
    cbind(method = "fg_smc", imps_fg_smc$impDatasets[[1]]),
    cbind(method = "fg_approx", complete(imps_cs_approx, action = 1L))
  return(imps)
})
```

5. Fit the Fine–Gray substantive model in each imputed dataset (using standard Cox software with I(D=1) and imputed V as outcome variables), and pool the estimates using Rubin's rules.

```
# Bind everything together
dat_imps <- rbindlist(list_imps, idcol = ".imp")
dat_imps</pre>
```

	.imp	method	id	time	D	X	Z	D1	D2
	<int></int>	<char></char>	<int></int>	<num></num>	<fctr></fctr>	<fctr></fctr>	<num></num>	<num></num>	<num></num>
1:	1	CCA	1	0.491195	0	1	0.126	0	0
2:	1	CCA	3	0.910797	0	0	-1.571	0	0
3:	1	CCA	8	0.036202	1	<na></na>	1.564	1	0
4:	1	CCA	9	0.046798	0	0	-1.653	0	0
5:	1	CCA	10	0.997413	0	<na></na>	-1.196	0	0
49996:	5	fg_approx	1992	0.319702	2	0	-2.670	0	1
49997:	5	fg_approx	1993	0.229071	2	0	-0.243	0	1
49998:	5	fg_approx	1994	1.836303	2	1	-0.366	0	1
49999:	5	fg_approx	1997	0.702380	2	0	0.283	0	1

```
50000:
                                        2
                                                  1 1.377
                                                              0
                                                                    1
           5 fg approx 1999 0.023554
               H1
                                                  Lambda1
                          H2 newtimes newevent
            <num>
                       <num>
                                <num>
                                        <fctr>
                                                     <num>
    1: 0.16736459 0.55436927 0.491195
                                             0 0.12385222
    2: 0.25761243 0.83833716 0.910797
                                             0 0.16659793
    3: 0.02028935 0.09603222 0.036202
                                             1 0.01932257
    4: 0.02606228 0.10990397 0.046798
                                             0 0.02452308
    5: 0.27549886 0.87116320 0.997413
                                             0 0.17340532
49996: 0.12370372 0.43826433 0.957205
                                             0 0.17116627
49997: 0.09740419 0.35023923 0.453168
                                             0 0.12098105
49998: 0.47538639 1.23075745 2.841599
                                             0.0.25988878
49999: 0.21877205 0.71087168 1.170590
                                             0 0.19454317
50000: 0.01356742 0.06584427 2.997529
                                             0 0.26284736
# To use the usual workflow: subset one of the methods first
imps fg smc <- dat imps[dat imps$method == "fg smc", ]</pre>
# Fit model in each imputed dataset
mods fg smc <- lapply(</pre>
 X = seq len(M),
 FUN = function(m) {
    imp_m <- imps_fg_smc[imps_fg_smc$.imp == m, ]</pre>
    coxph(Surv(newtimes, D1) ~ X + Z, data = imp m)
 }
)
# Pool results
summary(pool(mods fg smc))
 term estimate std.error statistic
                                              df
                                                      p.value
   X1 0.7768682 0.21722362 3.576352
                                        9.883541 5.136286e-03
     Z 0.4920664 0.06519244 7.547906 105.385333 1.659276e-11
# Alternative:
# Use (nested) {data.table} workflow to pool all methods simultaneously!
dat mods <- dat imps[, .(</pre>
 mod = list(coxph(Surv(newtimes, D1) ~ X + Z, data = .SD))
), by = c("method", ".imp")]
dat mods
      method
                            mod
              .imp
       <char> <int>
                         t>
 1:
          CCA
                1 <coxph[22]>
2:
       cs smc
                  1 <coxph[21]>
                  1 <coxph[21]>
3: cs approx
4:
                  1 <coxph[21]>
       fg smc
                  1 <coxph[21]>
5: fg_approx
                  2 <coxph[22]>
6:
         CCA
7:
                  2 <coxph[21]>
       cs smc
```

```
2 <coxph[21]>
 8: cs_approx
                   2 <coxph[21]>
 9:
       fg smc
                   2 <coxph[21]>
10: fg_approx
11:
           CCA
                   3 <coxph[22]>
12:
                   3 <coxph[21]>
       cs smc
                   3 <coxph[21]>
13: cs_approx
                   3 <coxph[21]>
14:
       fg_smc
                   3 <coxph[21]>
15: fg approx
                   4 <coxph[22]>
16:
           CCA
                   4 <coxph[21]>
17:
       \mathtt{cs\_smc}
18: cs_approx
                   4 <coxph[21]>
                   4 <coxph[21]>
19:
       fg smc
                   4 <coxph[21]>
20: fg_approx
21:
                   5 <coxph[22]>
           CCA
22:
                   5 <coxph[21]>
       cs smc
23: cs approx
                   5 <coxph[21]>
24:
       fg_smc
                   5 <coxph[21]>
25: fg_approx
                   5 <coxph[21]>
       method
                .imp
                              mod
```

#### dat\_mods[, summary(pool(as.list(mod))), by = "method"]

```
method
                term
                      estimate
                                 std.error statistic
                                                              df
                                                                       p.value
       <char> <fctr>
                          <niim>
                                     <num>
                                                <num>
                                                           <num>
                                                                         <num>
 1:
          CCA
                  X1 0.7781281 0.17916465
                                            4.343089 152.067624 2.554742e-05
 2:
          CCA
                   Z 0.4003856 0.10186017
                                            3.930737 145.744472 1.304356e-04
 3:
                  X1 0.6980657 0.18538543
                                            3.765483
                                                       14.973349 1.875994e-03
       cs smc
                                             7.769090
       cs_smc
                   Z 0.5079436 0.06538007
                                                       93.531830 9.965454e-12
 5: cs_approx
                  X1 0.6092265 0.19461615
                                            3.130400
                                                       12.205414 8.525728e-03
 6:
    cs_approx
                   Z 0.5225790 0.06779656
                                            7.708046
                                                       58.618467 1.775328e-10
                  X1 0.7768682 0.21722362
                                            3.576352
                                                        9.883541 5.136286e-03
 7:
       fg smc
 8:
       fg_smc
                   Z 0.4920664 0.06519244
                                            7.547906 105.385333 1.659276e-11
                  X1 0.6092265 0.19461615
 9: fg_approx
                                            3.130400
                                                       12.205414 8.525728e-03
10: fg_approx
                   Z 0.5225790 0.06779656
                                            7.708046
                                                       58.618467 1.775328e-10
```

# S2 Applied data example

# S2.1 Data dictionary

Table 1: Data dictionary. CMV: cytomegalovirus; HLA: human leukocyte antigen; HCT-CI: Hematopoietic stem cell transplantation-comorbidity index; MF: myelofibrosis.

Characteristic	N = 3,982
Patient age (years)	58 (52, 64)
Patient/donor CMV match	( ) /
Patient negative/Donor negative	1,142 (30%)
Other	2,715 (70%)
(Missing)	$1\overline{25}$
Donor type	
HLA identical sibling	1,183 (30%)
Other	2,795 (70%)
(Missing)	4
Hemoglobin (g/dL)	9.10 (8.10, 10.40)
(Missing)	1,873
HCT-CI risk category	
Low risk $(0)$	1,674 (54%)
Intermediate risk $(1-2)$	743~(24%)
High risk $(\geq 3)$	674~(22%)
(Missing)	891
Interval diagnosis-transplantation (years)	3(1, 9)
Karnosfky performance score	
$\geq 90$	2,475~(66%)
80	986~(26%)
$\leq 70$	$267 \ (7.2\%)$
(Missing)	254
Patient sex	
Female	1,484 (37%)
Male	2,498 (63%)
Peripheral blood (PB) blasts (%)	$1.0\ (0.0,\ 3.0)$
(Missing)	2,323
Conditioning	(
Standard	1,373 (35%)
Reduced	2,553~(65%)
(Missing)	56
Ruxolitinib given	
No	1,832 (66%)
Yes	931 (34%)
(Missing)	1,219
Disease subclassification	0.010 (50%)
Primary MF	2,912 (73%)
Secondary MF	1,070 (27%)
Night sweats	

No	1,256 (70%)
Yes	529 (30%)
(Missing)	$2,\!197$
T-cell depletion (in- or ev-vivo)	
No	1,012~(26%)
Yes	2,905 (74%)
(Missing)	65
Cytogenetics	
Normal	1,318 (59%)
Abnormal	910 (41%)
(Missing)	1,754
White blood cell count (WBC, $x10^9/L$ )	7(4, 14)
(Missing)	1,884
>10% Weight loss prior to transplantation	
No	1,329 (73%)
Yes	492~(27%)
(Missing)	2,161
Year of transplantation	2,015.0 (2,012.0, 2,018.0)

<sup>&</sup>lt;sup>1</sup> Median (IQR); n (%)

### S2.2 Non-parametric cumulative incidence curves

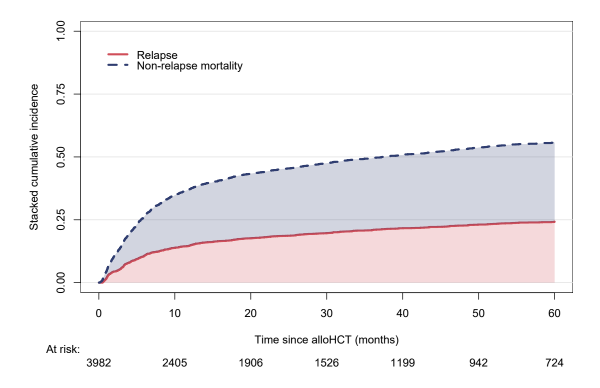


Figure 1: Stacked non-parametric cumulative incidence curves for competing relapse and non-relapse mortality, in dataset of 3982 primary and secondary myelofibrosis patients.

## S2.3 Pooled regression coefficients

Table 2: Pooled log hazard ratios [log HR, 95% confidence interval] for Fine–Gray model for relapse, cause-specific Cox model relapse, and cause-specific Cox model for non-relapse mortality (NRM).

Term + method	Relapse subdist. log HR	Relapse cause-spec. log HR	NRM cause-spec. log HR					
Conditioning: reduced								
CCA	0.02 [-0.33, 0.36]	0.01 [-0.33, 0.35]	0 [-0.29, 0.28]					
CS-SMC	0.13 [-0.02, 0.28]	0.1 [-0.05, 0.25]	-0.05 [-0.18, 0.07]					
CS-Approx	0.13 [-0.02, 0.28]	0.1 [-0.05, 0.25]	-0.05 [-0.18, 0.07]					
FG-SMC	0.13 [-0.02, 0.28]	0.1 [-0.05, 0.25]	-0.06 [-0.18, 0.07]					
FG-Approx	0.13 [-0.03, 0.28]	0.1 [-0.06, 0.25]	-0.05 [-0.18, 0.07]					
CMV match: o	ther							
CCA	0.04 [-0.31, 0.4]	0.05 [-0.3, 0.41]	0.09 [-0.19, 0.37]					
CS-SMC	-0.1 [-0.26, 0.05]	-0.05 [-0.2, 0.11]	0.22 [0.08, 0.36]					
CS-Approx	-0.1 [-0.26, 0.05]	-0.05 [-0.2, 0.11]	0.22 [0.08, 0.36]					
FG-SMC	-0.1 [-0.26, 0.05]	-0.04 [-0.2, 0.11]	0.22 [0.08, 0.36]					

(continued ...)

Table 2: (continued)

Term + method	Relapse subdist. log HR	Relapse cause-spec. log HR	NRM cause-spec. log HR
FG-Approx	-0.11 [-0.26, 0.05]	-0.05 [-0.2, 0.11]	$0.22 \ [0.08, \ 0.35]$
Cytogenetics: a	abnormal		
CCA	0.36 [0.04, 0.68]	0.37 [0.05, 0.68]	-0.08 [-0.35, 0.19]
CS-SMC	0.35 [0.15, 0.54]	0.35 [0.16, 0.54]	-0.07 [-0.23, 0.1]
CS-Approx	0.36 [0.17, 0.55]	0.35 [0.16, 0.54]	-0.08 [-0.25, 0.08]
FG-SMC	0.36 [0.17, 0.55]	0.36 [0.17, 0.54]	-0.06 [-0.21, 0.08]
FG-Approx	0.34 [0.17, 0.52]	0.34 [0.17, 0.51]	-0.07 [-0.22, 0.08]
Donor relation:	other		
CCA	0.12 [-0.28, 0.52]	0.2 [-0.2, 0.6]	0.53 [0.18, 0.88]
CS-SMC	-0.26 [-0.41, -0.1]	-0.19 [-0.34, -0.03]	0.35 [0.21, 0.5]
CS-Approx	-0.25 [-0.41, -0.1]	-0.18 [-0.34, -0.02]	0.36 [0.21, 0.5]
FG-SMC	-0.26 [-0.41, -0.1]	-0.19 [-0.34, -0.03]	0.35 [0.2, 0.49]
FG-Approx	-0.26 [-0.41, -0.1]	-0.19 [-0.34, -0.03]	0.35 [0.2, 0.49]
Hemoglobin (pe	er 5 g/dL)		
CCA	-0.38 [-0.85, 0.09]	-0.39 [-0.85, 0.08]	-0.12 [-0.49, 0.25]
CS-SMC	-0.24 [-0.51, 0.03]	-0.3 [-0.58, -0.03]	-0.19 [-0.42, 0.04]
CS-Approx	-0.25 [-0.53, 0.02]	-0.32 [-0.59, -0.06]	-0.19 [-0.41, 0.02]
FG-SMC	-0.25 [-0.51, 0.02]	-0.29 [-0.56, -0.02]	-0.08 [-0.28, 0.11]
FG-Approx	-0.23 [-0.5, 0.04]	-0.27 [-0.54, 0]	-0.09 [-0.29, 0.11]
<b>HCT-CI</b> $(1-2)$	)	, 1	, ,
CCA	-0.15 [-0.53, 0.22]	-0.04 [-0.42, 0.33]	0.38 [0.08, 0.69]
CS-SMC	-0.22 [-0.42, -0.01]	-0.17 [-0.37, 0.03]	0.15 [-0.02, 0.31]
CS-Approx	-0.19 [-0.38, 0.01]	-0.14 [-0.34, 0.06]	0.15 [-0.01, 0.31]
FG-SMC	-0.22 [-0.42, -0.01]	-0.18 [-0.38, 0.02]	0.12 [-0.04, 0.28]
FG-Approx	-0.19 [-0.38, 0.01]	-0.15 [-0.35, 0.04]	0.11 [-0.05, 0.27]
<b>HCT-CI</b> (≥ 3)			
CCA	-0.27 [-0.7, 0.16]	-0.19 [-0.62, 0.23]	0.4 [0.07, 0.73]
CS-SMC	-0.07 [-0.28, 0.14]	-0.01 [-0.21, 0.2]	0.27 [0.1, 0.44]
CS-Approx	-0.08 [-0.28, 0.13]	-0.02 [-0.22, 0.18]	0.26 [0.1, 0.43]
FG-SMC	-0.06 [-0.27, 0.14]	-0.02 [-0.22, 0.19]	0.21 [0.05, 0.37]
FG-Approx	-0.08 [-0.28, 0.11]	-0.04 [-0.23, 0.16]	0.21 [0.05, 0.38]
	sis to alloHCT (decades		ι , ,
CCA	0.01 [-0.24, 0.26]	0 [-0.25, 0.26]	-0.03 [-0.25, 0.19]
CS-SMC	-0.02 [-0.14, 0.09]	-0.02 [-0.14, 0.1]	0.05 [-0.05, 0.15]
CS-Approx	-0.03 [-0.14, 0.09]	-0.02 [-0.14, 0.1]	0.05 [-0.05, 0.15]
FG-SMC	-0.02 [-0.14, 0.09]	-0.02 [-0.13, 0.1]	0.05 [-0.05, 0.15]
FG-Approx	-0.02 [-0.14, 0.09]	-0.02 [-0.14, 0.1]	0.05 [-0.05, 0.15]
Karnofsky (80)	0.02 [ 0.11, 0.00]	0.02 [ 0.11, 0.1]	0.00 [ 0.00, 0.10]
CCA	-0.09 [-0.48, 0.31]	-0.08 [-0.48, 0.31]	0.04 [-0.27, 0.34]
CS-SMC	0.07 [-0.1, 0.24]	0.12 [-0.05, 0.28]	0.17 [0.03, 0.31]
CS-Approx	0.06 [-0.1, 0.24]	0.1 [-0.06, 0.27]	0.17 [0.03, 0.31]
FG-SMC	0.07 [-0.09, 0.24]	0.12 [-0.05, 0.29]	0.13 [0.01, 0.29] 0.17 [0.03, 0.31]
FG-Approx	0.07 [-0.03, 0.24]	0.12 [-0.06, 0.29]	0.17 [0.03, 0.31]
Karnofsky (≤ 7		0.12 [ 0.00, 0.23]	0.17 [0.00, 0.01]
CCA	0.63 [0.15, 1.11]	0.79 [0.3, 1.28]	0.33 [-0.13, 0.79]
CS-SMC	0.44 [0.19, 0.69]	0.79 [0.3, 1.28]	0.31 [0.08, 0.53]
CS-Approx	0.42 [0.17, 0.67]	0.51 [0.26, 0.76]	0.26 [0.04, 0.49]
FG-SMC	0.42 [0.17, 0.07] $0.44 [0.19, 0.7]$	0.55 [0.29, 0.81]	0.32 [0.09, 0.54]
FG-Approx	0.43 [0.17, 0.68]	0.53 [0.28, 0.78]	0.31 [0.08, 0.53]
		• • •	0.51 [0.06, 0.55]
CCA	sification: secondary M $-0.05$ [-0.45, 0.35]	-0.02 [-0.42, 0.38]	0.07 [-0.27, 0.41]
CS-SMC	0.01 [-0.17, 0.19]	0.01 [-0.17, 0.19]	0.07 [-0.27, 0.41] 0 [-0.16, 0.15]
CS-Approx	0 [-0.18, 0.18]	0 [-0.18, 0.19]	0 [-0.16, 0.15]
FG-SMC FG-Approx	0 [-0.18, 0.18] 0 [-0.18, 0.18]	0 [-0.18, 0.18] 0 [-0.18, 0.18]	-0.01 [-0.16, 0.15] -0.01 [-0.16, 0.15]
		0 [-0.18, 0.18]	-0.01 [-0.10, 0.15]
Night sweats: y	yes		
			/ 1

(continued ...)

Table 2: (continued)

Term + method	Relapse subdist. log HR	Relapse cause-spec. log HR	NRM cause-spec. log HR
CCA	-0.33 [-0.7, 0.04]	-0.4 [-0.77, -0.02]	-0.02 [-0.32, 0.27]
CS-SMC	-0.18 [-0.41, 0.05]	-0.2 [-0.44, 0.03]	-0.02 [-0.23, 0.19]
CS-Approx	-0.12 [-0.36, 0.13]	-0.14 [-0.38, 0.1]	0.03 [-0.19, 0.24]
FG-SMC	-0.17 [-0.4, 0.07]	-0.18 [-0.41, 0.05]	0.01 [-0.16, 0.19]
FG-Approx	-0.16 [-0.4, 0.07]	-0.18 [-0.42, 0.05]	0 [-0.17, 0.18]
Patient age (dec	cades)		
CCA	0.1 [-0.09, 0.28]	0.13 [-0.06, 0.32]	0.13 [-0.02, 0.28]
CS-SMC	-0.03 [-0.12, 0.05]	0.01 [-0.08, 0.09]	$0.21 \ [0.14, \ 0.29]$
CS-Approx	-0.03 [-0.12, 0.05]	0.01 [-0.08, 0.09]	$0.21 \ [0.14, \ 0.29]$
FG-SMC	-0.04 [-0.12, 0.05]	0.01 [-0.08, 0.09]	0.22 [0.15, 0.3]
FG-Approx	-0.03 [-0.12, 0.05]	0.01 [-0.08, 0.09]	0.22 [0.15, 0.3]
Patient sex: ma		0.40 [ 0.84   0.48]	
CCA	-0.24 [-0.56, 0.09]	-0.18 [-0.51, 0.15]	0.39 [0.11, 0.68]
CS-SMC	-0.1 [-0.24, 0.05]	-0.06 [-0.21, 0.09]	0.18 [0.05, 0.31]
CS-Approx	-0.1 [-0.24, 0.05]	-0.06 [-0.21, 0.09]	0.18 [0.05, 0.31]
FG-SMC FG-Approx	-0.09 [-0.24, 0.05] -0.1 [-0.24, 0.05]	-0.06 [-0.2, 0.09] -0.06 [-0.21, 0.08]	0.18 [0.05, 0.31] 0.18 [0.05, 0.31]
PB Blasts (per	= = = = = = = = = = = = = = = = = = = =	-0.00 [-0.21, 0.06]	0.10 [0.00, 0.01]
CCA	0.16 [-0.04, 0.36]	0.17 [-0.02, 0.37]	0 [-0.18, 0.18]
CS-SMC	0.18 [0.05, 0.31]	0.17 [-0.02, 0.37]	0.01 [-0.12, 0.13]
CS-Approx	0.19 [0.07, 0.31]	0.18 [0.03, 0.31] $0.19 [0.07, 0.32]$	0.01 [-0.12, 0.13]
FG-SMC	0.17 [0.04, 0.3]	0.17 [0.05, 0.3]	-0.01 [-0.12, 0.13]
FG-Approx	0.18 [0.05, 0.32]	0.18 [0.05, 0.31]	-0.02 [-0.12, 0.09]
Ruxolitinib give		L / 1	, ,
CCA	0.08 [-0.26, 0.43]	0.08 [-0.26, 0.43]	-0.05 [-0.33, 0.23]
CS-SMC	-0.02 [-0.2, 0.17]	-0.03 [-0.22, 0.16]	-0.06 [-0.21, 0.1]
CS-Approx	$0.01 \left[ -0.19, 0.2 \right]$	-0.01 [-0.2, 0.18]	-0.05 [-0.21, 0.11]
FG-SMC	-0.02 [-0.21, 0.17]	-0.03 [-0.22, 0.16]	-0.04 [-0.19, 0.11]
FG-Approx	0 [-0.19, 0.18]	-0.01 [-0.2, 0.17]	-0.04 [-0.19, 0.11]
T-cell depletion			
CCA	0.2 [-0.21, 0.62]	0.16 [-0.25, 0.58]	-0.23 [-0.54, 0.08]
CS-SMC	0.3 [0.13, 0.48]	0.26 [0.09, 0.44]	-0.18 [-0.32, -0.04]
CS-Approx	0.3 [0.12, 0.48]	0.26 [0.08, 0.43]	-0.19 [-0.33, -0.05]
FG-SMC	0.31 [0.13, 0.48]	0.26 [0.09, 0.44]	-0.18 [-0.31, -0.04]
FG-Approx	0.31 [0.13, 0.48]	$0.26 \ [0.09, \ 0.44]$	-0.18 [-0.32, -0.04]
WBC count (log			
CCA	0.17 [0.02, 0.33]	0.17 [0.01, 0.33]	0.02 [-0.12, 0.15]
CS-SMC	0.17 [0.09, 0.26]	0.18 [0.09, 0.27]	0 [-0.07, 0.07]
CS-Approx	0.17 [0.08, 0.26]	0.17 [0.09, 0.26]	0 [-0.08, 0.07]
FG-SMC FG-Approx	0.17 [0.09, 0.26] 0.17 [0.1, 0.25]	0.18 [0.09, 0.26] 0.18 [0.1, 0.26]	-0.01 [-0.07, 0.05] -0.01 [-0.08, 0.05]
	• • •	0.18 [0.1, 0.20]	-0.01 [-0.06, 0.05]
Weight loss: yes CCA	0 [-0.37, 0.38]	0.05 [-0.33, 0.43]	0.17 [ 0.12   0.48]
CS-SMC	0.23 [-0.03, 0.49]	0.03 [-0.53, 0.45]	0.17 [-0.13, 0.48] 0.16 [-0.05, 0.36]
CS-Approx	0.24 [0, 0.47]	0.28 [0.04, 0.51]	0.16 [-0.05, 0.36]
FG-SMC	0.23 [-0.01, 0.47]	0.24 [0.01, 0.48]	0.06 [-0.12, 0.24]
FG-Approx	0.24 [0, 0.48]	0.24 [0.01, 0.40] $0.26 [0.02, 0.49]$	0.06 [-0.12, 0.24]
Year of alloHC7	• • •	ŗ , <u>-</u> 1	r , -1
CCA	-0.36 [-0.99, 0.26]	-0.41 [-1.04, 0.23]	-0.15 [-0.67, 0.37]
CS-SMC	-0.08 [-0.34, 0.18]	-0.11 [-0.37, 0.15]	-0.24 [-0.46, -0.02]
CS-Approx	-0.09 [-0.35, 0.17]	-0.12 [-0.38, 0.14]	-0.24 [-0.46, -0.02]
FG-SMC	-0.08 [-0.34, 0.17]	-0.12 [-0.37, 0.14]	-0.24 [-0.46, -0.03]
FG-Approx	-0.08 [-0.34, 0.17]	-0.11 [-0.37, 0.14]	-0.24 [-0.46, -0.03]