

iCARE (Individualized Coherent Absolute Risk Estimation) Package

December 10, 2018

Load the iCARE library

```
> library(iCARE)
```

Load the breast cancer data and set the seed.

```
> data("bc_data", package="iCARE")
> set.seed(50)
```

Example 1: SNP-only model

In this example, we will estimate the risk of breast cancer in ages 50-80. A SNP-only model is fit, with no specific genotypes supplied for estimation. The population disease rates are from SEER.

```
> res_snps_miss = computeAbsoluteRisk(model.snp.info = bc_15_snps,
+                                     model.disease.incidence.rates = bc_inc,
+                                     model.competing.incidence.rates = mort_inc,
+                                     apply.age.start = 50, apply.age.interval.length = 30,
+                                     return.refs.risk = TRUE)

Note: You did not provide apply.snp.profile. Will impute SNPs for 10000 people.
If require more, please provide apply.snp.profile input.
[1] "Note: As specified, the model does not adjust SNP imputations for family history."
    user   system elapsed
 19.296   0.458  26.315
```

Compute a summary of the risks.

```
> summary(res_snps_miss$refs.risk)
```

Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
0.07652	0.09198	0.09575	0.09583	0.09950	0.11966

Next, suppose we want to predict risk for three specific women whom we have genotyped; we can then call:

```

> res_snps_dat = computeAbsoluteRisk(model.snp.info = bc_15_snps,
+                                     model.disease.incidence.rates = bc_inc,
+                                     model.competing.incidence.rates = mort_inc,
+                                     apply.age.start = 50, apply.age.interval.length = 30,
+                                     apply.snp.profile = new_snp_prof,
+                                     return.refs.risk = TRUE)

[1] "Note: As specified, the model does not adjust SNP imputations for family history."
    user   system elapsed
0.559   0.267   1.465

> names(res_snps_dat)
[1] "risk"      "details"    "beta.used"  "refs.risk"

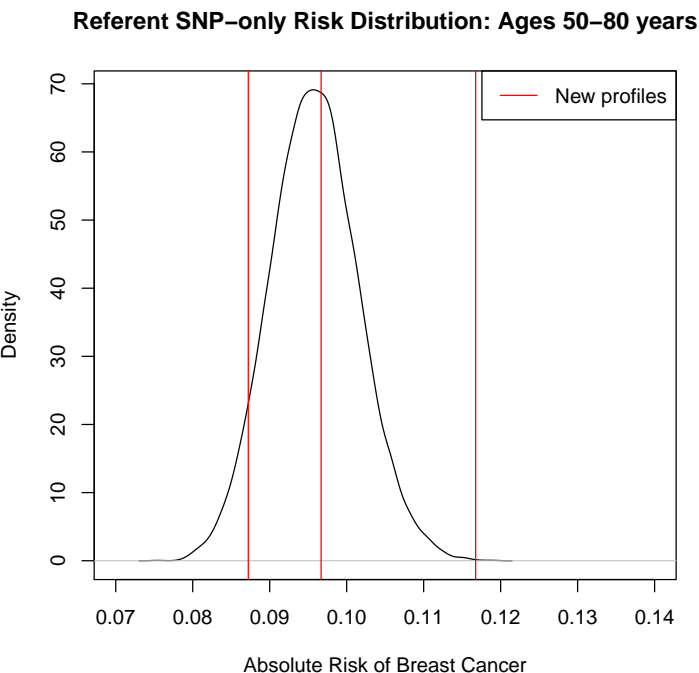
```

These results allow us to create a useful plot showing the distribution of risks in our reference dataset and to add the risks of the three women to see where they fall on the population distribution.

```

> plot(density(res_snps_dat$refs.risk),
+       xlim = c(0.07,0.14), xlab = "Absolute Risk of Breast Cancer",
+       main = "Referent SNP-only Risk Distribution: Ages 50-80 years")
> abline(v = res_snps_dat$risk, col = "red")
> legend("topright", legend = "New profiles", col = "red", lwd = 1)

```



Example 2: Breast cancer risk model with risk-factors and SNPs

In this example, we will estimate the risk of breast cancer in ages 50-80 by fitting a model with two risk factors and 15 SNPs, with three specific covariate profiles supplied for estimation (with some missing data). First, the model.cov.info argument is created.

```
> v1 = list(); v1$name = "famhist"; v1$type = "continuous"
> v2 = list(); v2$name = "parity"; v2$type = "factor";
> v2$levels = c(0,1,2,3,4)
> bc_model_cov_info = list(v1,v2)
> bc_model_formula = observed.outcome ~ famhist + as.factor(parity)
> res_covs_snps = computeAbsoluteRisk(model.formula = bc_model_formula,
+                                         model.cov.info = bc_model_cov_info,
+                                         model.snp.info = bc_15_snps,
+                                         model.log.RR = bc_model_log_or,
+                                         model.ref.dataset = ref_cov_dat,
+                                         model.disease.incidence.rates = bc_inc,
+                                         model.competing.incidence.rates = mort_inc,
+                                         model.bin.fh.name = "famhist",
+                                         apply.age.start = 50,
+                                         apply.age.interval.length = 30,
+                                         apply.cov.profile = new_cov_prof,
+                                         apply.snp.profile = new_snp_prof,
+                                         return.refs.risk = TRUE)

      user  system elapsed
0.093   0.038   0.194
```

In addition to summarizing and plotting the risk estimates, iCARE includes an option to view more detailed output, by calling:

```
> print(res_covs_snps$details)
```

	Int_Start	Int_End	Risk_Estimate	rs12405132	rs12048493	rs72755295	
193502	50	80	0.08077312	NA	NA	0	
126252	50	80	0.07709955	0	0	0	
15756	50	80	0.12955390	0	1	0	
	rs6796502	rs13162653	rs2012709	rs7707921	rs9257408	rs4593472	rs13365225
193502	0	1	1	0	1	1	1
126252	0	2	1	0	1	1	0
15756	1	0	1	0	1	1	0
	rs13267382	rs11627032	rs146699004	rs745570	rs6507583	famhist	parity
193502	0	0	1	2	0	0	2
126252	1	1	0	1	0	0	4
15756	1	0	0	0	0	1	4

Illustration of the validation component

We want to validate a model for predicting absolute risk of disease based on a combined model of two epidemiologic risk factors family history and parity and

15 SNPs using the nested case-control dataset.

The first step is to compute sampling weights. We fit a logistic regression model of inclusion depending on the case/control status, age of study entry and observed followup using the R function **glm**, as shown below:

```
> validation.cohort.data$inclusion = 0
> subjects_included = intersect(validation.cohort.data$id,
+                                 validation.nested.case.control.data$id)
> validation.cohort.data$inclusion[subjects_included] = 1
> validation.cohort.data$observed.followup =
+   validation.cohort.data$study.exit.age -
+   validation.cohort.data$study.entry.age
> selection.model = glm(inclusion ~ observed.outcome
+                         * (study.entry.age + observed.followup),
+                         data = validation.cohort.data,
+                         family = binomial(link = "logit"))
> validation.nested.case.control.data$sampling.weights =
+   selection.model$fitted.values[validation.cohort.data$inclusion == 1]
```

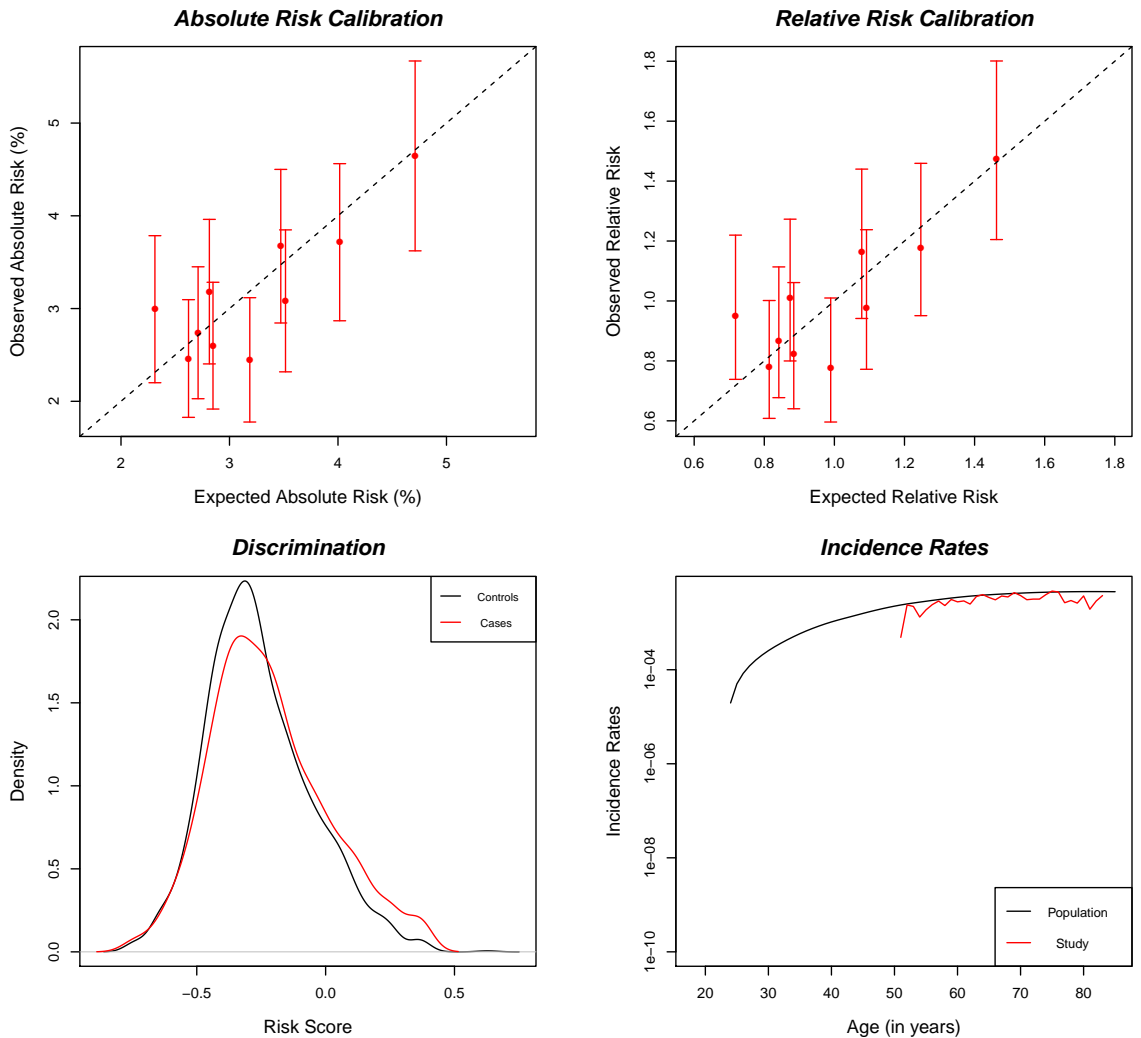
The next step is to call the **ModelValidation** function to implement the validation analysis.

```
> bc_model_formula = observed.outcome ~ famhist + as.factor(parity)
> data = validation.nested.case.control.data
> risk.model = list(model.formula = bc_model_formula,
+                     model.cov.info = bc_model_cov_info,
+                     model.snp.info = bc_15_snps,
+                     model.log.RR = bc_model_log_or,
+                     model.ref.dataset = ref_cov_dat,
+                     model.ref.dataset.weights = NULL,
+                     model.disease.incidence.rates = bc_inc,
+                     model.competing.incidence.rates = mort_inc,
+                     model.bin.fh.name = "famhist",
+                     apply.cov.profile = data[,all.vars(bc_model_formula)[-1]],
+                     apply.snp.profile = data[,bc_15_snps$snp.name],
+                     n.imp = 5, use.c.code = 1, return.lp = TRUE,
+                     return.refs.risk = TRUE)
> output = ModelValidation(study.data = data,
+                           total.followup.validation = TRUE,
+                           predicted.risk.interval = NULL,
+                           iCARE.model.object = risk.model,
+                           number.of.percentiles = 10)

      user  system elapsed
3.878    0.012   3.940
```

We can also produce a set of useful plots showing the results of the validation analysis.

```
> plotModelValidation(study.data = data, validation.results = output)
NULL
```



Dataset: Example Dataset

Model Name: Example Model

Risk Prediction Interval: Observed Followup

Number of subjects (cases): 2694 (1112)

Follow-up time (years) [mean,range]: [9.782 , (5 , 13)]

Baseline age (years) [mean,range]: [62.721 , (50 , 72)]

E/O [Estimate, 95% CI]: [1.024 , (0.948 , 1.107)]

Absolute Risk Calibration

HL Test, df: 11.086 , 10
p-value: 3.50876e-01

Relative Risk Calibration

Test, df: 10.685 , 9
p-value: 2.979384e-01

Model Discrimination

AUC est: 0.54
95% CI: (0.517 , 0.563)

Session Information

```
> sessionInfo()

R version 3.5.1 Patched (2018-07-12 r74967)
Platform: x86_64-apple-darwin15.6.0 (64-bit)
Running under: OS X El Capitan 10.11.6

Matrix products: default
BLAS: /Library/Frameworks/R.framework/Versions/3.5/Resources/lib/libRblas.0.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/3.5/Resources/lib/libRlapack.dylib

locale:
[1] C/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8

attached base packages:
[1] stats      graphics   grDevices utils      datasets   methods    base

other attached packages:
[1] iCARE_1.10.3    Hmisc_4.1-1     ggplot2_3.1.0   Formula_1.2-3
[5] survival_2.43-3 lattice_0.20-38 gtools_3.8.1   plotrix_3.7-4

loaded via a namespace (and not attached):
[1] Rcpp_1.0.0          pillar_1.3.0       compiler_3.5.1
[4] RColorBrewer_1.1-2  plyr_1.8.4        bindr_0.1.1
[7] base64enc_0.1-3     tools_3.5.1       digest_0.6.18
[10] rpart_4.1-13       checkmate_1.8.5  htmlTable_1.12
[13] tibble_1.4.2        gtable_0.2.0      pkgconfig_2.0.2
[16] rlang_0.3.0.1       Matrix_1.2-15    rstudioapi_0.8
[19] bindr_0.2.2         gridExtra_2.3    stringr_1.3.1
[22] knitr_1.20          withr_2.1.2      dplyr_0.7.8
[25] cluster_2.0.7-1    htmlwidgets_1.3   grid_3.5.1
[28] nnet_7.3-12         tidyselect_0.2.5  data.table_1.11.8
[31] glue_1.3.0          R6_2.3.0         foreign_0.8-71
[34] latticeExtra_0.6-28 purrr_0.2.5     magrittr_1.5
[37] htmltools_0.3.6     backports_1.1.2  scales_1.0.0
[40] splines_3.5.1       assertthat_0.2.0  colorspace_1.3-2
[43] stringi_1.2.4       acepack_1.4.1    lazyeval_0.2.1
[46] munsell_0.5.0       crayon_1.3.4
```