

## Complex sampling and R.

**Thomas Lumley** 

**UW** Biostatistics

useR, Rennes — 2009-7-7

## Why are surveys special?

- Probability samples come with a lot of meta-data that has to be linked correctly to the observations, so software was specialized
- Interesting data sets tend to be large, so hardware was specialized
- Design-based inference literature is largely separate from rest of statistics, doesn't seem to have a unifying concept such as likelihood to rationalize arbitrary choices.

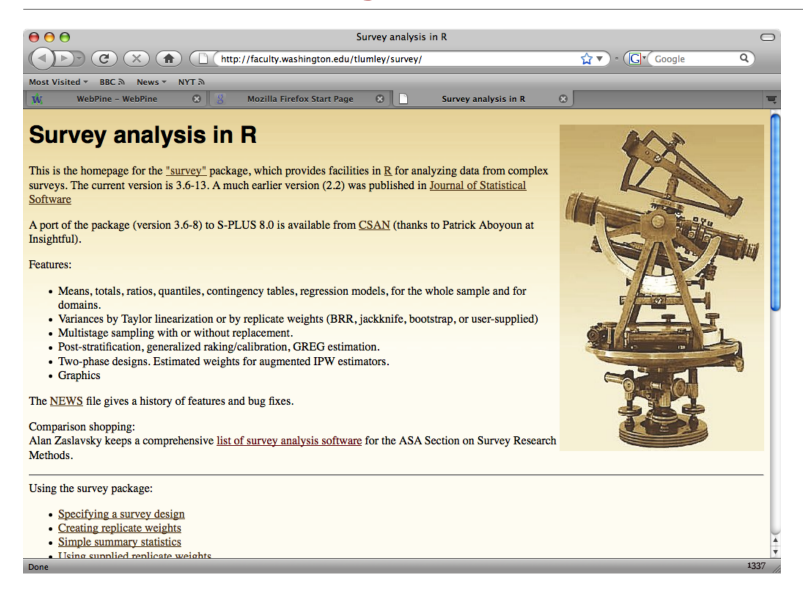
In part, too, the specialized terminology has formed barriers around the territory. At one recent conference, I heard a speaker describe methods used in his survey with the sentence, We used BRR on a PPS sample of PSUs, with MI after NRFU

—Sharon Lohr, The American Statistician, 5/2004

## Times are changing

- Connections between design-based inference and semiparametric models
- Increasing use of 'sandwich' standard errors in biostatistics
- Connections between sampling weights and causal inference
- Increasing availability of design-based methods in standard software
- Increasing use of complex designs in epidemiology

## R Survey package



http://faculty.washington.edu/tlumley/survey/

## R Survey package

Version 3.16 is current, containing approximately 9000 lines of interpreted R code. (cf 250,000 lines of Fortran for VPLX)

Version 2.3 was published in Journal of Statistical Software.

Major changes since then are finite population corrections for multistage sampling, calibration and generalized raking, tests of independence in contingency tables, better tables of results, better graphics, loglinear models, ordinal regression models, two-phase designs, database-backed designs, PPS designs.

## Design principles

- Ease of maintenance and debugging by code reuse
- Speed and memory use not initially a priority: dont optimize until there are real use-cases to profile.
- Rapid release, so that bugs and other infelicities can be found and fixed.
- Emphasize features that look like biostatistics (regression, calibration, survival analysis, exploratory graphics)

### Intended market

- Methods research (because of programming features of R)
- Teaching (because of cost, use of R in other courses)
- Secondary analysis of national surveys (regression features,
   R is familiar to non-survey statisticians)
- Two-phase designs in epidemiology

#### **Outline**

- Describing survey designs: svydesign()
- Database-backed designs
- Summary statistics: mean, total, quantiles, design effect
- Tables of summary statistics, domain estimation.
- Graphics: histograms, hexbin scatterplots, smoothers.
- Regression modelling: svyglm()
- Multiply-imputed data
- Calibration of weights: postStratify(), calibrate()

## **Objects and Formulas**

Collections of related information should be kept together in an object. For surveys this means the data and the survey metadata.

The way to specify variables from a data frame or object in R is a formula

$$^{a} + b + I(c < 5*d)$$

The survey package always uses formulas to specify variables in a survey data set.

### **Basic estimation ideas**

Individuals are sampled with known probabilities  $\pi_i$  from a population of size N to end up with a sample of size n. The population can be a full population or an existing sample such as a cohort.

We write  $R_i = 1$  if individual i is sampled,  $R_i = 0$  otherwise

The design-based inference problem is to estimate what any statistic of interest would be if data from the whole population were available.

### **Basic estimation ideas**

For a population total this is easy: an unbiased estimator of

$$T_X = \sum_{i=1}^N x_i$$

is

$$\widehat{T}_X = \sum_{i:R_i=1} \frac{1}{\pi_i} X_i$$

Standard errors follow from formulas for the variance of a sum: main complication is that we do need to know  $cov[R_i, R_j]$ .

### **Basic estimation ideas**

Other statistics follow from sums: if the statistic on the whole population would solve the estimating equation

$$\sum_{i=1}^{N} U_i(\theta) = 0$$

then a design-based estimate will solve

$$\sum_{i:R_i=1} \frac{1}{\pi_i} U_i(\theta) = 0$$

Standard errors come from the delta method or from resampling (actually reweighting).

Theoretical details can become tricky, especially if  $U_i()$  is not a function of just one observation.

Stratified independent sample (without replacement) of California schools

- stype is a factor variable for elementary/middle/high school
- fpc is a numeric variable giving the number of schools in each stratum. If omitted we assume sampling with replacement
- $id=\sim 1$  specifies independent sampling.
- apistrat is the data frame with all the data.
- pw contains sampling weights  $(1/\pi_i)$ . These could be omitted since they can be computed from the population size.

Note that all the variables are in apistrat and are specified as formulas.

```
> dstrat
Stratified Independent Sampling design
svydesign(id = ~1, strata = ~stype, weights = ~pw, data = apistrat,
fpc = fpc
> summary(dstrat)
Stratified Independent Sampling design
svydesign(id = ~1, strata = ~stype, weights = ~pw, data = apistrat,
fpc = fpc
Probabilities:
Min. 1st Qu. Median Mean 3rd Qu. Max.
0.02262 0.02262 0.03587 0.04014 0.05339 0.06623
Stratum Sizes:
F. H M
obs 100 50 50
design.PSU 100 50 50
actual.PSU 100 50 50
Population stratum sizes (PSUs):
F. M H
4421 1018 755
Data variables:
[1] "cds" "stype" "name" "sname" "snum" "dname"
[7] "dnum" "cname" "cnum" "flag" "pcttest" "api00"
. . .
```

Cluster sample of school districts, using all schools within a district.

```
dclus1 <- svydesign(id=~dnum, weights=~pw, data=apiclus1, fpc=~fpc)</pre>
```

- dnum is a (numeric) identifier for school district
- No stratification, so no strata= argument

```
> summary(dclus1)
1 - level Cluster Sampling design
With (15) clusters.
svydesign(id = ~dnum, weights = ~pw, data = apiclus1, fpc = ~fpc)
Probabilities:
Min. 1st Qu. Median Mean 3rd Qu. Max.
0.02954 0.02954 0.02954 0.02954 0.02954
Population size (PSUs): 757
Data variables:
[1] "cds" "stype" "name" "sname" "snum" "dname"
[7] "dnum" "cname" "cnum" "flag" "pcttest" "api00"
...
```

Two-stage sample: 40 school districts and up to 5 schools from each

```
dclus2 <- svydesign(id=~dnum+snum, fpc=~fpc1+fpc2, data=apiclus2)</pre>
```

- dnum identifies school district, snum identifies school
- fpc1 is the number of school districts in population, fpc2 is number of schools in the district.
- Weights are computed from fpc1 and fpc2

```
> summary(dclus2)
2 - level Cluster Sampling design
With (40, 126) clusters.
svydesign(id = ~dnum + snum, fpc = ~fpc1 + fpc2, data = apiclus2)
Probabilities:
Min. 1st Qu. Median Mean 3rd Qu. Max.
0.003669 0.037740 0.052840 0.042390 0.052840 0.052840
Population size (PSUs): 757
Data variables:
[1] "cds" "stype" "name" "sname" "snum" "dname"
[7] "dnum" "cname" "cnum" "flag" "pcttest" "api00"
...
```

California Health Interview Survey has 50 sets of replicate weights instead of design information

Can also create replicate weights with as.svrepdesign(), using jackknife, bootstrap, or BRR.

## Describing surveys: database-based

```
library(RSQLite)
brfss <- svydesign(id="X_PSU, strata="X_STATE, weight="X_FINALWT,
    data="brfss", dbtype="SQLite", dbname="brfss07.db",
    nest=TRUE)</pre>
```

The data argument is the name of a database table or view, dbtype and dbname specify the database.

Only the design meta-data are loaded into the design object. Other variables are temporarily loaded as needed when an analysis is run.

Can use any database with an ODBC, JDBC, or R-DBI interface: anything on Windows, SQLite, Postgres, Oracle.

BRFSS is about as large as a 1Gb laptop can handle with this approach: 430,000 records.

## **Summary statistics**

svymean, svytotal, svyratio, svyvar, svyquantile

All take a formula and design object as arguments, return an object with coef, vcov, SE, cv methods.

Mean and total on factor variables give tables of cell means/totals.

Mean and total have deff argument for design effects and the returned object has a deff method.

```
> svymean(~api00, dclus1, deff=TRUE)
mean SE DEff
api00 644.169 23.542 9.3459
> svymean(~factor(stype),dclus1)
mean SE
factor(stype)E 0.786885 0.0463
factor(stype)H 0.076503 0.0268
factor(stype)M 0.136612 0.0296
```

## **Summary statistics**

```
> svymean(~interaction(stype, comp.imp), dclus1)
mean SE
interaction(stype, comp.imp)E.No 0.174863 0.0260
interaction(stype, comp.imp)H.No 0.038251 0.0161
interaction(stype, comp.imp)M.No 0.060109 0.0246
interaction(stype, comp.imp)E.Yes 0.612022 0.0417
interaction(stype, comp.imp)H.Yes 0.038251 0.0161
interaction(stype, comp.imp)M.Yes 0.076503 0.0217
> svyvar(~api00, dclus1)
variance SE
api00 11183 1386.4
> svytotal(~enroll, dclus1, deff=TRUE)
total SE DEff
enroll 3404940 932235 31.311
```

## **Summary statistics**

```
> mns <- svymean(~api00+api99,dclus1)</pre>
> mns
      mean SE
api00 644.17 23.542
api99 606.98 24.225
> coef(mns)
api00 api99
644.1694 606.9781
> SE(mns)
api00 api99
23.54224 24.22504
> vcov(mns)
     api00 api99
api00 554.2371 565.7856
api99 565.7856 586.8526
> cv(mns)
api00 api99
0.03654666 0.03991090
```

### **Domain estimation**

The correct standard error estimate for a subpopulation that isnt a stratum is not just obtained by pretending that the subpopulation was a designed survey of its own.

However, the subset function and "[" method for survey design objects handle all these details automagically, so you can ignore this problem.

The package test suite (tests/domain.R) verifies that subpopulation means agree with derivations from ratio estimators and regression estimator derivations. Some more documentation is in the domain vignette.

Note: subsets of design objects don't necessarily use less memory than the whole objects.

### **Prettier tables**

Two main types

• totals or proportions cross-classified by multiple factors

• arbitrary statistics in subgroups

svyby computes a statistic for subgroups specified by a set of factor variables:

```
> svyby(~api99, ~stype, dclus1, svymean)
stype statistics.api99 se.api99
E E 607.7917 22.81660
H H 595.7143 41.76400
M M 608.6000 32.56064
```

~api99 is the variable to be analysed, ~stype is the subgroup variable, dclus1 is the design object, svymean is the statistic to compute.

Lots of options for eg what variance summaries to present

```
> svyby(~api99, ~stype, dclus1, svyquantile, quantiles=0.5,ci=TRUE)
  stype statistics.quantiles
                                 statistics.CIs
                                                      se
                                                               var
F.
      Ε
                         615 525.6174, 674.1479 37.89113 1435.738
Η
      Η
                         593 428.4810, 701.0065 69.52309
М
     M
                         611 527.5797, 675.2395 37.66903 1418.955
> svyby(~api99, list(school.type=apiclus1$stype), dclus1, svymean)
  school.type statistics.api99 se.api99
Ε
                      607.7917 22.81660
Η
            Η
                      595.7143 41.76400
M
                      608.6000 32.56064
> svyby(~api99+api00, ~stype, dclus1, svymean, deff=TRUE)
> svyby(~api99+api00, ~stype, dclus1, svymean, deff=TRUE)
  stype statistics.api99 statistics.api00 se.api99 se.api00
F.
      F.
                607.7917
                                 648.8681 22.81660 22.36241
Η
                595.7143
                                 618.5714 41.76400 38.02025
M
                608,6000
                                 631.4400 32.56064 31.60947
 DEff.api99 DEff.api00
F.
    5.895734 6.583674
H 2.211866 2.228259
    2.226990 2.163900
```

	stype	sch.wide	statistic.api99	statistic.api00
E.No	E	No	601.6667	596.3333
H.No	H	No	662.0000	659.3333
M.No	M	No	611.3750	606.3750
E.Yes	Ε	Yes	608.3485	653.6439
H.Yes	H	Yes	577.6364	607.4545
M.Yes	M	Yes	607.2941	643.2353

```
> (a<-svyby(~enroll, ~stype, rclus1, svytotal, deff=TRUE,
+ vartype=c("se","cv","cvpct","var")))
  stype statistics.enroll
                               se cv.enroll cv%.enroll
                                                                          DEff
                                                                var
               2109717.1 631349.4 0.2992578
Ε
                                              29.92578 398602047550 125.039075
Η
                535594.9 226716.6 0.4232987 42.32987 51400414315
                                                                      4.645816
М
                759628.1 213635.5 0.2812369
                                              28.12369 45640120138 13.014932
> deff(a)
[1] 125.039075 4.645816 13.014932
> SE(a)
[1] 631349.4 226716.6 213635.5
> cv(a)
[1] 0.2992578 0.4232987 0.2812369
> coef(a)
[1] 2109717.1 535594.9 759628.1
> svyby(~api00,~comp.imp+sch.wide,design=dclus1,svymean,
+ drop.empty.groups=FALSE)
        comp.imp sch.wide statistics.api00 se.api00
No.No
             No
                      No
                                 608.0435 28.98769
Yes.No
            Yes
                      No
                                       NΑ
                                                NA
No.Yes
             No
                     Yes
                                 654.0741 32.66871
Yes. Yes
                  Yes
                               648,4060 22,47502
            Yes
```

### **Functions of estimates**

svycontrast computes linear and nonlinear combinations of estimated statistics (in the same object).

### **Functions of estimates**

### **Crosstabs**

svyby or svymean and svytotal with interaction will produce the numbers, but the formatting is not pretty.

#### ftable provides formatting:

```
> d<-svyby(~api99 + api00, ~stype + sch.wide, rclus1, svymean,
         keep.var=TRUE, vartype=c("se","cvpct"))
> round(ftable(d),1)
           sch.wide
                              No
                                                                Yes
             statistics.api99 statistics.api00 statistics.api99 statistics.api00
stype
      svymean
                        601.7
                                         596.3
                                                           608.3
                                                                             653.6
Ε
      SE
                         70.0
                                          64.5
                                                            23.7
                                                                              22.4
      cv%
                         11.6
                                          10.8
                                                             3.9
                                                                               3.4
                        662.0
                                         659.3
                                                           577.6
                                                                             607.5
Η
      svymean
                         40.9
                                          37.8
                                                            57.4
                                                                              54.0
      SE
      cv%
                          6.2
                                                             9.9
                                           5.7
                                                                               8.9
                       611.4
                                         606.4
                                                           607.3
                                                                             643.2
M
      svymean
                         48.2
      SE
                                          48.3
                                                            49.5
                                                                              49.3
      cv%
                          7.9
                                           8.0
                                                             8.2
                                                                               7.7
```

#### **Crosstabs**

svyby knows enough to structure the table without help. For other analyses more information is needed

```
> a<-svymean(~interaction(stype,comp.imp), design=dclus1, deff=TRUE)</pre>
> b<-ftable(a, rownames=list(stype=c("E","H","M"),comp.imp=c("No","Yes")))</pre>
> round(100*b,1)
                        E
                              Η
                                     M
              stype
comp.imp
                     17.5
No
                            3.8
                                 6.0
         mean
         SE
                      2.6
                            1.6
                                 2.5
         Deff
                     87.8 131.7 200.4
                     61.2
Yes
                             3.8
                                   7.7
         mean
         SE
                      4.2
                            1.6
                                   2.2
         Deff
                    137.2 131.4 124.7
```

## Testing in tables

svychisq does four variations on the Pearson  $\chi^2$  test: corrections to the mean or mean and variance of  $X^2$  (Rao and Scott) and two Wald-type tests (Koch et al).

The exact asymptotic distribution of the Rao–Scott tests (linear combination of  $\chi_1^2$ ) is also available.

## Loglinear models

svyloglin() does loglinear models

```
a<-svyloglin(~backpain+neckpain+sex+sickleave, nhis)
a2<-update(a, ~.^2)
a3<-update(a, ~.^3)
anova(a, a2)
anova(a2, a3)
b1<-update(a, ~.+(backpain*neckpain*sex)+sex*sickleave)
b2<-update(a, ~.+(backpain+neckpain)*sex+sex*sickleave)
b3<-update(a, ~.+backpain:neckpain+sex:backpain+sex:neckpain
+sex:sickleave)</pre>
```

anova() method computes Rao-Scott working loglikelihood and working score tests, with the two Rao-Scott approximations for the p-value and the exact asymptotic distribution.

## **Graphics**

When complex sampling designs are analyzed with regression models it is more important to have good exploratory analysis methods.

Problems with survey data

- large data
- unequal weights.

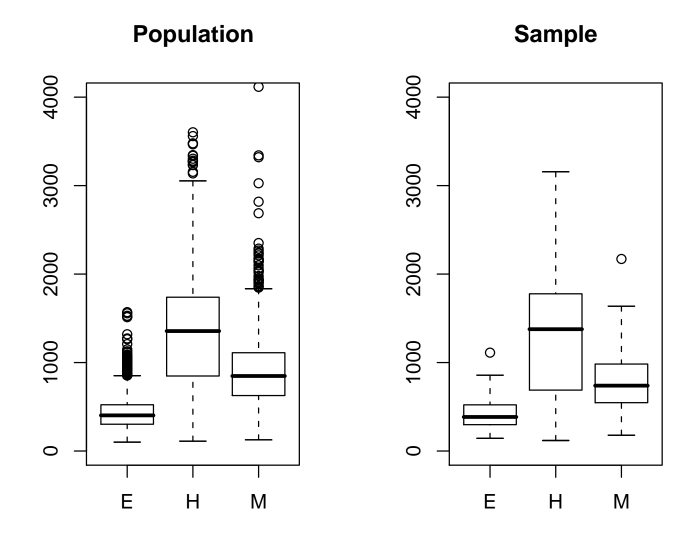
## **Estimating populations**

Boxplots, barplots, and histograms display estimates of the population distribution function for a variable.

We can substitute the survey estimates: boxplots use quantiles, histograms and barplots need tables. eg boxplot of enrollment by school type (Elementary/Middle/High)

svyboxplot(enroll~stype, design=srs)

## **Estimating populations**



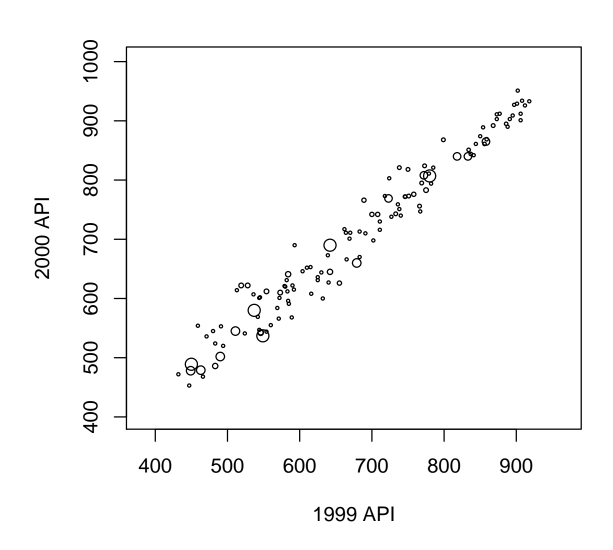
This approach doesn't work for scatterplots. We need to incorporate weights in the plotting symbols.

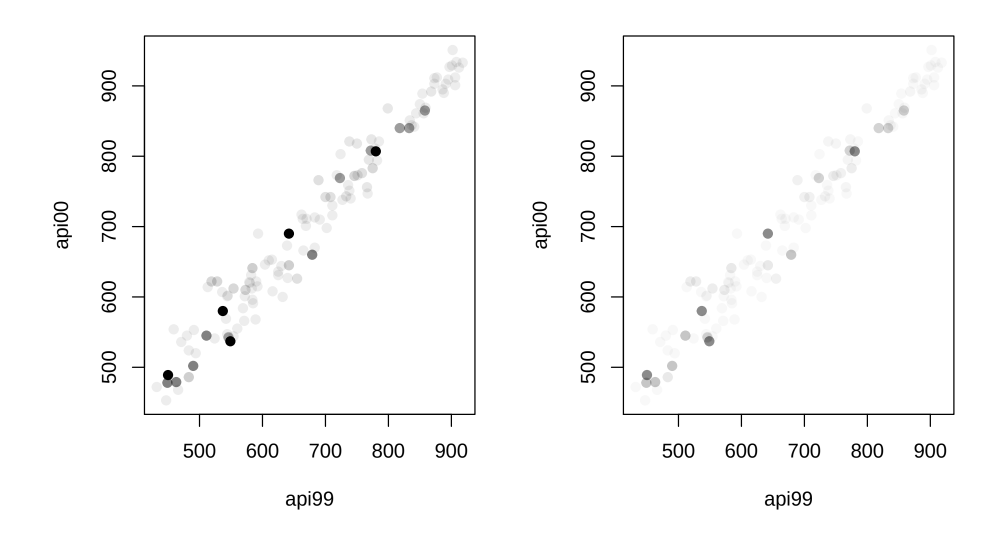
One approach is the bubble plot: radius or area of circle proportional to sampling weight.

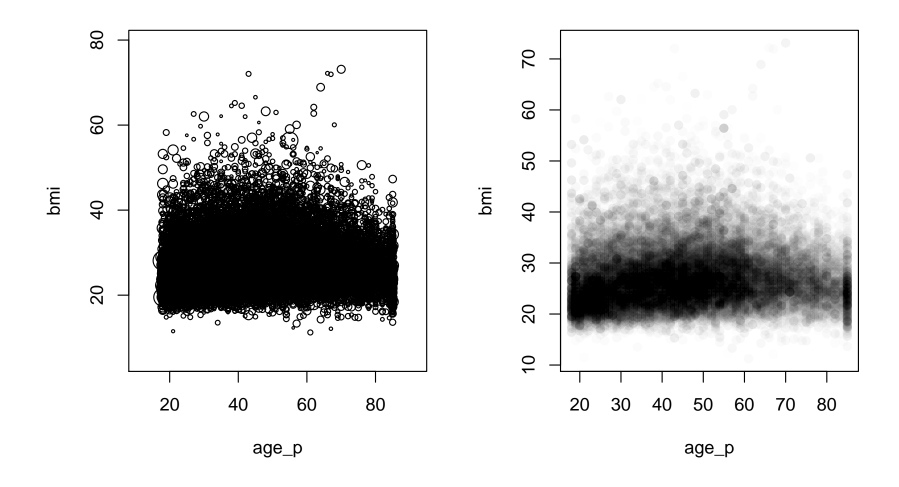
Another is transparent color: opacity for each point is proportional to sampling weight, giving a density estimation effect.

Bubble plots work well with small data sets, badly with large data sets.

Transparency works well with large data sets, badly with small data sets.







### Binning and smoothing

Hexagonal binning divides the plotting area into hexagons, counts the number in each hexagon [Carr, 1987, JASA]. In dense regions this has similar effect to transparency, but allows outliers to be seen, and produces much smaller graphics files.

For survey data we just add up the weight in each bin to get the estimated population numbers in each bin.

In R: svyplot(bmi~age\_p, design=nhis, style="hex")}

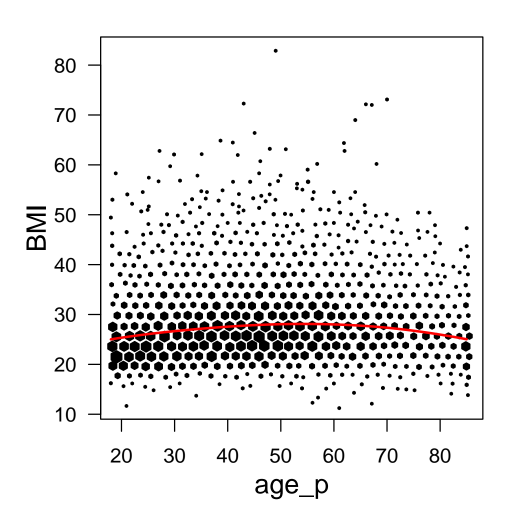
### Binning and smoothing

Scatterplot smoothers fit a smooth curve through the middle of the data. There are many variants, but they all work by estimating the mean value of Y using points in a small interval of values of x. svysmooth() uses a fast binned kernel smoother (Matt Wand).

In R use svysmooth to create a smooth curve and then plot to draw it.

```
smth <- svysmooth(api00~api99, stratrs, bandwidth=40)
plot(smth)</pre>
```

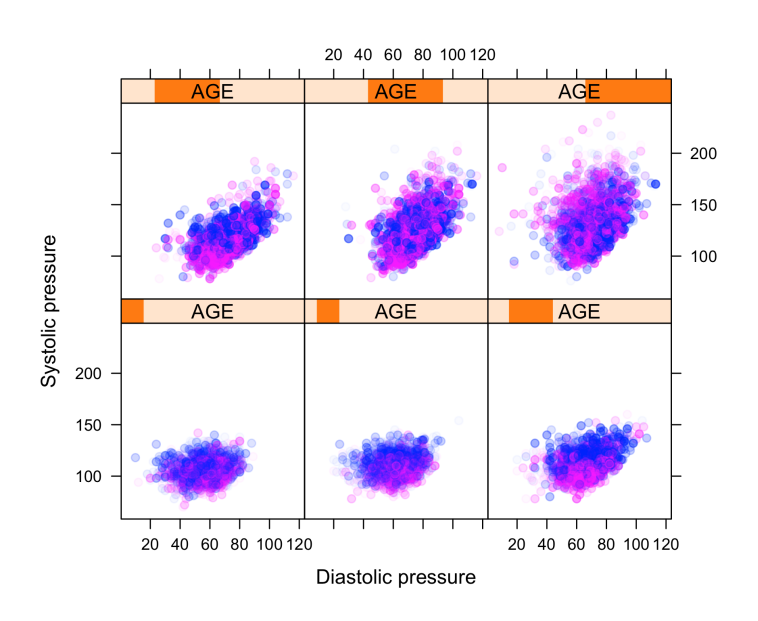
### Binning and smoothing



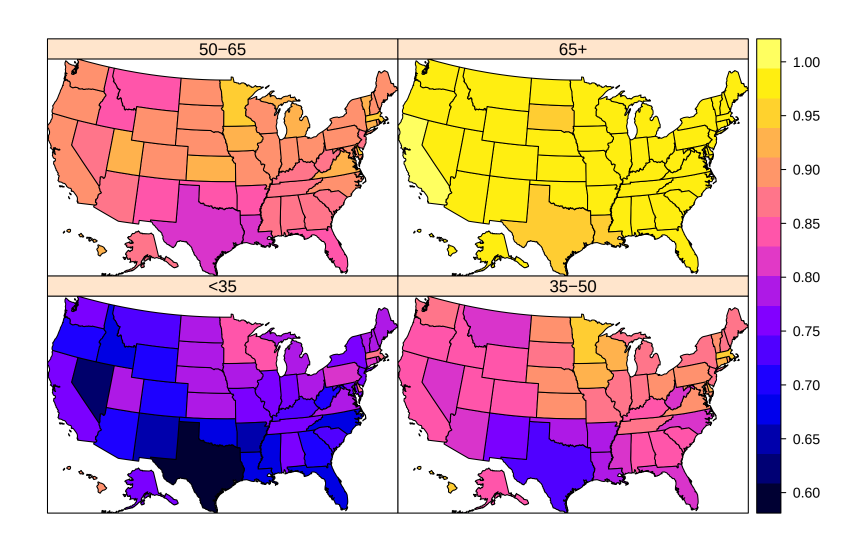
#### Counts

- **2265009**
- **1**2123446
- **1981883**
- **1840320**
- 1698757
- **1557194**
- **1415631**
- 1274068
- 1132505
- 990942
- 849379
- 707816
- 566253
- 424690
- 283127
- 141564
- •

# Blood pressure by age and gender



# Health insurance by age and state



```
svycoplot(sysbp~diabp|agegp, style="transparent",
   basecol=function(d) c("magenta", "royalblue") [d$sex]
   data=nhanes_design)
library(maptools)
states<-readShapePoly("brfss_state_2007_download")</pre>
states<-states[order(states$ST_FIPS),]</pre>
brfss<-update(brfss, agegp=cut(AGE, c(0,35,50,65,Inf)))
hlth<-svyby(~I(HLTHPLAN==1), ~agegp+X_STATE, svymean,
    design=brfss)
hlthdata<-reshape(hlth[,c(1,2,4)],idvar="X_STATE",
    direction="wide",timevar="agegp")
names(hlthdata)[2:5] <-paste("age",1:4,sep="")</pre>
states@data<-merge(states,hlthdata,
    by.x="ST_FIPS",by.y="X_STATE",all=FALSE)
spplot(states,c("age1","age2","age3","age4"),
    names.attr=c("<35","35-50","50-65","65+"))
```

### Regression models

- svyglm for linear and generalized linear models
- svyolr for proportional odds and other cumulative link models.
- svycoxph for Cox model (no std errors on survival curves yet)

For these models the point estimates are the same for frequency weights, sampling weights, or precision weights, so the point estimation can just reuse the ordinary regression code.

### Regression estimator of total

Simple random sample of 200 counties in US. Use 2000 election results to help predict 2008 totals

```
m.obama<-svyglm(OBAMA~BUSH+GORE, srsdes)</pre>
m.mccain<-svyglm(MCCAIN~BUSH+GORE, srsdes)</pre>
data2000<-data.frame(BUSH=sum(elections$BUSH),</pre>
        GORE=sum(elections$GORE))
predict(m.obama, total=3049, newdata=data2000)
predict(m.mccain, total=3049, newdata=data2000)
1.mccain<-svyglm(MCCAIN~BUSH+GORE, srsdes,</pre>
    family=quasi(variance="mu"))
1.obama<-svyglm(OBAMA~BUSH+GORE, srsdes,</pre>
    family=quasi(variance="mu"))
predict(1.obama, total=3049, newdata=data2000)
predict(1.mccain, total=3049, newdata=data2000)
```

### Regression estimator of total

HT estimator: standard error 29 million for Obama, 13 million for McCain

Regression estimator: std error 6 million for Obama, 4 million for McCain

Variance proportional to mean: std error 4.1 million for Obama, 3.7 million for McCain.

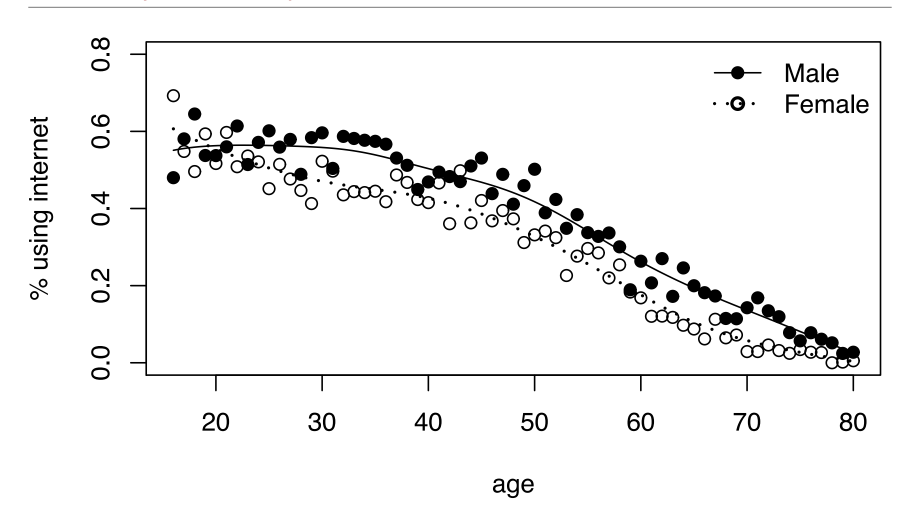
Precision gain is a free lunch: no assumptions needed for validity.

Regression can also help with non-response bias (but does make assumptions). Take Indiana and Virginia rather than a random sample.

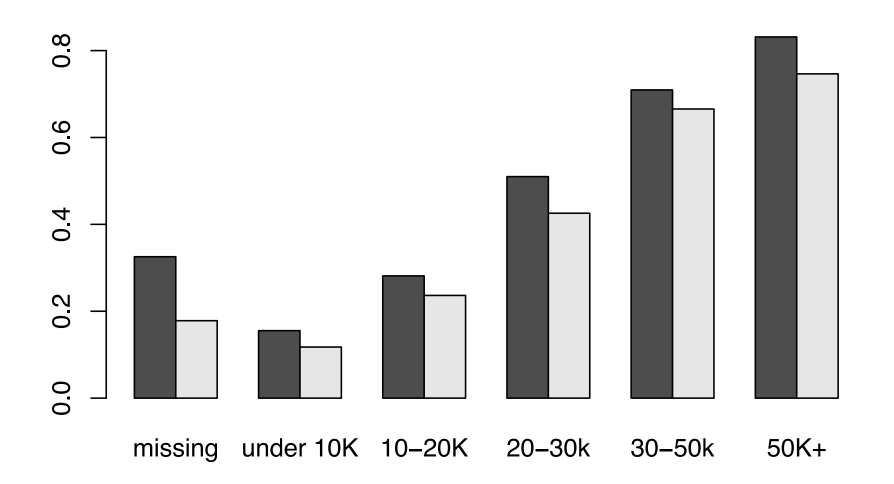
### Regression model

Internet use in Scotland (2001) by age, sex, and income. shs<-svydesign(id=~psu, strata=~stratum, weight=~grosswt, data=shs data) bys<-svyby(~intuse,~age+sex,svymean,design=shs)</pre> plot(svysmooth(intuse~age, design=subset(shs,sex=="male" & !is.na(age)), bandwidth=5),ylim=c(0,0.8),ylab="% using internet") lines(svysmooth(intuse~age, design=subset(shs,sex=="female" & !is.na(age)), bandwidth=5),lwd=2,lty=3) points(bys\$age,bys\$intuse,pch=ifelse(bys\$sex=="male",19,1)) legend("topright",pch=c(19,1),lty=c(1,3),lwd=c(1,2), legend=c("Male", "Female"), bty="n") byinc<-svyby(~intuse, ~sex+groupinc, design=shs)</pre> barplot(byinc)

# Age (cohort) effect



### **Income**



```
> m<-svyglm(intuse~I(age-18)*sex,design=shs,
   family=quasibinomial())
> m2<-svyglm(intuse~(pmin(age,35)+pmax(age,35))*sex,
   design=shs,family=quasibinomial)
> summary(m)
svyglm(intuse ~ I(age - 18) * sex, design = shs,
  family = quasibinomial())
Survey design:
svydesign(id = ~psu, strata = ~stratum, weight = ~grosswt,
  data = ex2)
Coefficients:
                      Estimate Std. Error t value Pr(>|t|)
(Intercept)
                      0.804113 0.047571 16.903 < 2e-16 ***
I(age - 18)
                    -0.044970 0.001382 -32.551 < 2e-16 ***
sexfemale
                    -0.116442 0.061748 -1.886 0.0594.
I(age - 18):sexfemale -0.010145
                                0.001864 -5.444 5.33e-08 ***
```

```
> summary(m2)
Call:
svyglm(intuse ~ (pmin(age, 35) + pmax(age, 35)) * sex,
   design = shs, family = quasibinomial)
Survey design:
svydesign(id = ~psu, strata = ~stratum, weight = ~grosswt,
  data = ex2
Coefficients:
                        Estimate Std. Error t value Pr(>|t|)
(Intercept)
                        2.152291 0.156772 13.729 < 2e-16 ***
pmin(age, 35)
                       0.014055
                                  0.005456 2.576 0.010003 *
                      -0.063366
pmax(age, 35)
                                  0.001925 - 32.922 < 2e-16 ***
sexfemale
                        0.606718
                                  0.211516 2.868 0.004133 **
                                  0.007294 -2.352 0.018691 *
pmin(age, 35):sexfemale -0.017155
pmax(age, 35):sexfemale -0.009804
                                  0.002587 -3.790 0.000151 ***
```

### Extending the package

More general models eg censored parametric regression models, can be fitted with svymle, which takes a loglikelihood and a set of model formulas for one or more of the parameters in the likelihood.

If code is available to get point estimates (eg using precision weights), withReplicates() will produce replicate-weight standard errors

svymle is still limited to models where the objective function is a sum of single-observation loglikelihoods: no mixed models.

### Example: negative binomial regression

Negative binomial regression is not implemented in the survey package, but it is implemented for simple random samples in the MASS package

Model for mean

$$\log E[Y] = \log \mu_i = \alpha + X_i \beta \tag{1}$$

and variance

$$\operatorname{var}\left[Y\right] = \mu_i + \mu_i^2/\theta. \tag{2}$$

The likelihood for one observation is

$$L(\mu_i, \theta) = \frac{\Gamma(Y_i + \theta)}{\Gamma(\theta)\Gamma(Y_i + 1)} \left(\frac{\theta}{\theta + \mu_i}\right)^{\theta} \left(\frac{\mu_i}{\mu_i + \theta}\right)^{Y_i}$$

#### **Options**

- 1. Fit a loglinear Poisson model instead, using svyglm()
- 2. For a replicate-weights design, rely on glm.nb for point estimates and use withReplicates() to estimate standard errors.
- 3. Use svymle().

#### **Example**

Data on lifetime number of sexual partners, collected by NHANES 2003–2004, and how it varies with age, gender, and race/ethnicity.

### Poisson regression

Fit Poisson regression with svyglm(), and compare the point estimates to glm.nb() library(MASS) des <- update(des, scaledweights = WTINT2YR/mean(WTINT2YR))</pre> model0 <- glm.nb(</pre> total factor (RIAGENDR) \* (log(age) + factor (RIDRETH1)), data=model.frame(des)) model1 <- glm.nb(</pre> total factor (RIAGENDR) \* (log(age) + factor (RIDRETH1)), data=model.frame(des), weights=scaledweights) model2 <- svyglm(</pre>

total factor (RIAGENDR) \* (log(age) + factor (RIDRETH1)),

design=des, family=quasipoisson)

### Poisson regression

```
> round(cbind(coef(model0), coef(model1), coef(model2)), 2)
                                  [,1] [,2] [,3]
(Intercept)
                                    2.35 2.29 2.44
                                   -0.91 -0.80 -0.95
factor(RIAGENDR)2
log(age)
                                    1.06 1.07 0.80
factor(RIDRETH1)2
                                    0.03 0.08 0.05
                                    0.07 0.09 0.06
factor(RIDRETH1)3
                                    1.00 0.82 0.71
factor(RIDRETH1)4
factor(RIDRETH1)5
                                   -0.01 0.06 0.01
factor(RIAGENDR)2:log(age)
                                   -1.26 -1.22 -0.96
factor(RIAGENDR)2:factor(RIDRETH1)2 0.04 -0.18 -0.16
factor(RIAGENDR)2:factor(RIDRETH1)3 0.68 0.60 0.63
factor(RIAGENDR)2:factor(RIDRETH1)4 -0.03 0.06 0.18
factor(RIAGENDR)2:factor(RIDRETH1)5 0.64 0.38 0.41
```

#### Replicate weights

glm.nb() in the MASS package fits negative binomial regression and accepts weights as an argument.

Looking at the code shows that the weights are used to multiply individual contributions to the loglikelihood, so that the point estimates will be correct if the weights are sampling weights.

```
repldesign <- as.svrepdesign(des)
negbin <- withReplicates(repldesign,
    quote(coef(glm.nb(
        total~factor(RIAGENDR)*(log(age)+factor(RIDRETH1)),
        weights=.weights))))</pre>
```

#### Replicate weights

Using starting values for the iteration should be faster.

```
model <- glm.nb(
    total~factor(RIAGENDR)*(log(age)+factor(RIDRETH1)),
    weights=weights(des), data=model.frame(des))
init.coef<-coef(model)
init.theta<-model$theta
negbin2 <- withReplicates(repldesign,
    quote(coef(glm.nb(
    total~factor(RIAGENDR)*(log(age)+factor(RIDRETH1)),
    weights=.weights, init.theta=init.theta, start=init.coef))))</pre>
```

svymle() will maximize a sampling-weighted estimate of a population loglikelihood or other additive objective function.

$$\ell_i = \log \Gamma(Y_i + \theta) - \log \Gamma(\theta) - \log \Gamma(Y_i + 1) + \theta \log \theta + Y_i \log(\mu_i) - (\theta + Y_i) \log(\theta + \mu_i)$$

$$\frac{\partial \ell}{\partial \mu_i} = \frac{Y_i}{\mu_i} - \frac{\theta + Y_i}{\theta + \mu_i}$$

$$\frac{\partial \ell}{\partial \theta} = \psi(\theta + Y_i) - \psi(\theta) + \log(\theta) + 1 - \log(\theta + \mu_i) - \frac{Y_i + \theta}{\mu_i + \theta}.$$

 $\psi$  is the digamma function, digamma() in R.

We will be specifying a linear model for  $\eta=\log\mu$ , so the derivative with respect to  $\mu$  needs to be multiplied by  $\partial\mu/\partial\eta=\mu$ 

```
loglik <- function(y, theta, eta) {</pre>
    mu<-exp(eta)</pre>
    (lgamma(theta + y) - lgamma(theta) - lgamma(y + 1) +
      theta * log(theta) + y * log(mu + (y == 0))
      - (theta + y) * log(theta + mu))
deta<- function(y, theta, eta) {</pre>
    mu <- exp(eta)</pre>
    dmu <- y/mu - (theta+y)/(theta+mu)</pre>
    dmu*mu
```

```
dtheta <- function(y, theta, eta) {</pre>
    mu <- exp(eta)</pre>
    digamma(theta + y) - digamma(theta) + log(theta) + 1
       - log(theta + mu) - (y + theta)/(mu + theta)
  }
score<-function(y, theta, eta) {</pre>
     cbind(dtheta(y, theta,eta), deta(y, theta, eta))
}
modl1<-glm.nb(</pre>
    total~factor(RIAGENDR)*(log(age)+factor(RIDRETH1)),
    data=model.frame(des), weights=scaledweights)
```

#### Results

```
> round(cbind(coef(nlmfit), SE(nlmfit), c(NA, coef(negbin)),
    c(NA, SE(negbin))),2)
                                         [,1] [,2] [,3] [,4]
theta.(Intercept)
                                         0.81 0.30
                                                      NA
                                                          NΑ
                                         2.29 0.16 2.29 0.16
eta.(Intercept)
eta.factor(RIAGENDR)2
                                        -0.80 0.18 -0.80 0.18
eta.log(age)
                                         1.07 0.23 1.07 0.24
                                         0.08 0.15 0.08 0.15
eta.factor(RIDRETH1)2
eta.factor(RIDRETH1)3
                                         0.09 0.18 0.09 0.18
                                         0.82 0.30 0.82 0.30
eta.factor(RIDRETH1)4
eta.factor(RIDRETH1)5
                                         0.06 0.38 0.06 0.41
eta.factor(RIAGENDR)2:log(age)
                                        -1.22 \ 0.26 \ -1.22 \ 0.27
eta.factor(RIAGENDR)2:factor(RIDRETH1)2 -0.18 0.26 -0.18 0.26
eta.factor(RIAGENDR)2:factor(RIDRETH1)3 0.60 0.19 0.60 0.20
eta.factor(RIAGENDR)2:factor(RIDRETH1)4 0.06 0.37 0.06 0.38
eta.factor(RIAGENDR)2:factor(RIDRETH1)5 0.38 0.44 0.38 0.46
```

#### **Simulations**

An advantage of using R for survey analysis is easier simulations.

A case—control study could be analyzed as a stratified random sample (stratified on case status), as survey statisticians do, or by maximum likelihood, as biostatisticians do.

We can compare these analyses by simulation [cf Scott & Wild, JRSSB 2000].

```
population<-data.frame(x=rnorm(10000))
expit <- function(eta) exp(eta)/(1+exp(eta))
population$y<- rbinom(10000, 1, expit(population$x-4))
population$wt<-ifelse(population$y==1, 1,(10000-sum(population$y))/600)

one.sample<-function(){
   cases<-which(population$y==1)
   controls<-sample(which(population$y==0),600)
   population[c(cases,controls),]
}</pre>
```

#### **Simulations**

```
mle<-function(sample) coef(glm(y~x, data=sample, family=binomial()))
svy<-function(sample){
   design<-svydesign(id=~1,strat=~y, weight=~wt, data=sample)
   coef(svyglm(y~x, design=design, family=quasibinomial()))
}</pre>
```

Testing shows that it seems to work, and 3 replicates takes about 1 second. 1000 replicates will take about 5 minutes.

#### **Simulations**

The true parameter values are (-4,1). Based on 1000 simulations the bias and variability of the estimators are

So the MLE has a relative efficiency of nearly 2 for the slope estimate and is biased for the intercept, as expected.

### Misspecified model

Try the analysis when the true model is not linear

$$logit P[Y = 1 | X = x] = e^x - 4$$

but still fit

$$logitP[Y = 1 | X = x] = \alpha + \beta x$$

'True' answer fitting logistic model to population is (-3.66, 0.75)

MLE is biased for population result but has lower variance.

data(esoph) in R are data from a case—control study of esophageal cancer in Ille-et-Vilaine. The study examined smoking and alcohol consumption as risk factors.

Alcohol consumption is coded into categories 0-39, 40-79, 80-119, 120+ grams/day, smoking into 0-9, 10-19, 20-29, 30+ grams/day.

Age is an extremely strong risk factor, and so is quite likely to be a confounder. It is coded in 10-year categories.

```
> summary(model1)
Call:
glm(formula = cbind(ncases, ncontrols) ~ agegp + tobgp + alcgp,
   family = binomial(), data = esoph)
Deviance Residuals:
   Min
             10
                 Median
                             3Q
                                     Max
-1.6891 -0.5618 -0.2168
                          0.2314
                                  2.0643
Coefficients:
           Estimate Std. Error z value Pr(>|z|)
(Intercept) -5.9108
                       1.0302 -5.737 9.61e-09 ***
agegp35-44
             1.6095
                       1.0676 1.508 0.131652
agegp45-54 2.9752
                       1.0242 2.905 0.003675 **
                       1.0198 3.293 0.000991 ***
agegp55-64
           3.3584
agegp65-74
            3.7270
                       1.0253 3.635 0.000278 ***
agegp75+
             3.6818
                       1.0645 3.459 0.000543 ***
tobgp10-19
            0.3407
                       0.2054 1.659 0.097159 .
tobgp20-29
            0.3962
                       0.2456 1.613 0.106708
tobgp30+
            0.8677
                       0.2765 3.138 0.001701 **
alcgp40-79
            1.1216
                       0.2384 4.704 2.55e-06 ***
alcgp80-119
             1.4471
                       0.2628 5.506 3.68e-08 ***
alcgp120+
             2.1154
                       0.2876 7.356 1.90e-13 ***
```

Alcohol and tobacco both show increasing association with increasing dose. We need tests for the set of coefficients for each of these variables.

Likelihood ratio tests are natural for maximum likelihood

Wald tests can be extended more easily to probability-weighted models

```
> library(survey)
> regTermTest(model1,~alcgp)
Wald test for alcgp
  in glm(formula = cbind(ncases, ncontrols) ~ agegp + tobgp + alcgp,
     family = binomial(), data = esoph)
Chisq = 57.89887 on 3 df: p= 1.652e-12
> regTermTest(model1,~tobgp)
Wald test for tobgp
  in glm(formula = cbind(ncases, ncontrols) ~ agegp + tobgp + alcgp,
     family = binomial(), data = esoph)
Chisq = 10.76880 on 3 df: p= 0.013044
```

We do not have the sampling fractions for controls, but the incidence of esophageal cancer is about 5/100,000 per year. We do not know how many years of cases were recruited, but it is reasonable to assume the sampling fraction is no smaller than 1/2000 for controls.

Using the survey package we can fit the same model with probability weights. It is necessary first to expand the data to one record per person, then to declare the sampling design, then use svyglm to fit the model.

The estimated odds ratios based on a sampling fraction of 1/2000 are very similar to the unweighted analysis, and the intercept is different, as expected.

```
> desoph<-svydesign(id=~1, weight=~prob, data=expanded)</pre>
> pmodel1<-svyglm(cancer~agegp+tobgp+alcgp, design=desoph, family=binomial)
> round(rbind(unweighted=coef(model1), weighted=coef(pmodel1)),3)
          (Intercept) agegp35-44 agegp45-54 agegp55-64 agegp65-74 agegp75+
              -5.911
                          1.610
                                    2.975
                                              3.358
                                                         3.727
                                                                 3,682
unweighted
weighted
                          1.607
                                              3.321
                                                         3.644
                                                                 3.657
              -13.422
                                    2.995
          tobgp10-19 tobgp20-29 tobgp30+ alcgp40-79 alcgp80-119 alcgp120+
unweighted
              0.341
                         0.396
                                 0.868
                                            1.122
                                                       1.447
                                                                2.115
weighted
              0.275
                         0.340 0.782
                                            1.104
                                                       1.437
                                                                2.004
```

In this example the standard errors are also similar

> round(rbind(unweighted=SE(model1), weighted=SE(pmodel1)),2)

```
(Intercept) agegp35-44 agegp45-54 agegp55-64 agegp65-74 agegp75+
                            1.07
                                       1.02
unweighted
                 1.03
                                                  1.02
                                                            1.03
                                                                     1.06
weighted
                 1.01
                            1.06
                                       1.02
                                                  1.01
                                                            1.02
                                                                     1.06
          tobgp10-19 tobgp20-29 tobgp30+ alcgp40-79 alcgp80-119 alcgp120+
                0.21
                           0.25
                                    0.28
                                              0.24
                                                          0.26
unweighted
                                                                    0.29
                                    0.29
                                              0.24
weighted
                0.21
                           0.25
                                                          0.27
                                                                    0.30
```

Sometimes the probability-weighted analysis is very inefficient, sometimes it is reasonably efficient.

### Two-phase sampling

- ullet Phase 1: sample people according to some probability design  $\pi_{1,i}$ , measure variables
- ullet Phase 2: subsample people from the phase 1 sample using the variables measured at phase 1,  $\pi_{2|1,i}$  and measure more variables

Sampling probability  $\pi_i = \pi_{1,i} \times \pi_{2|1,i}$  and use  $1/\pi_i$  as sampling weights.

These are **not** (in general) the marginal probability that i is in the sample, because  $\pi_{2|1,i}$  may depend on which other observations are in the phase-one sample.

### Two-phase sampling

For all the designs today  $\pi_{1,i}$  is constant, so its value doesn't affect anything except estimated population totals.

We don't care about population totals in this setting, so we don't need to know the value of  $\pi_{1,i}$  (eg can set to 1).

An alternative view is that we are using model-based inference at phase one, rather than sampling-based inference.

### Two-phase sampling

Two sources of uncertainty:

- Phase-1 sample is only part of population
- Phase 2 observes full set of variables on only a subset of people.

Uncertainties at the two phases add.

# Designs (survey)

The classic survey example is 'two-phase sampling for stratification'. This is useful when a good stratifying variable is not available for the population but is easy to measure.

- Take a large simple random sample or cluster sample and measure stratum variables
- Take a stratified random sample from phase 1 for the survey

If the gain from stratification is larger than the cost of phase 1 we have won.

# Designs (epi)

Many large cohorts exist in epidemiology. These can be modelled as simple random samples. They have a lot of variables measured.

It is common to want to measure a new variable

- New assay on stored blood
- Coding of open-text questionnaire
- Re-interview

The classic designs are a simple random sample and a case—control sample.

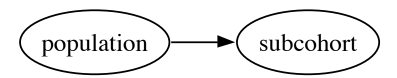
It is often more useful to sample based on multiple phase-1 variables: outcome, confounders, surrogates for phase-2 variable.

# **Analysis**

The true sampling is two-phase



We can ignore the first phase and pretend that we had an unstratified population sample



This is conservative, but sometimes not very. It was used before software for eg Cox models in two-phase samples were developed.

### Two-phase case-control

The case—control design stratifies on Y. We can stratify on X as well

	X=0	X=1	
Y=0	а	b	$m_0$
Y=1	С	d	$m_1$
	$n_{0}$	$n_1$	

The estimated variance of  $\beta$  is

$$\operatorname{var}[\widehat{\beta}] = \frac{1}{a} + \frac{1}{b} + \frac{1}{c} + \frac{1}{d}$$

Ideally want all cells about the same size in this trivial case.

### **Example: Wilm's Tumor**

- Wilm's Tumor is a rare childhood cancer of the kidney.
   Prognosis is good in early stage or favourable histology disease.
- Histology is difficult to determine. NWTSG central pathologist is much better than anyone else.
- To reduce cost of followup, consider central histology only for a subset of cases.
- Sample all relapses, all patients with unfavorable histology by local pathologist, 10% of remainder

## **Analyses: sampling weights**

#### Phase 1

```
relapse
instit 0 1
good 3207 415
bad 250 156
```

#### Phase 2

```
relapse
instit 0 1
good 314 415
bad 250 156
```

Weight is 1/sampling fraction in relapse  $\times$  local histology strata: 1.0 for cases, controls with unfavorable local histology, 3207/314 = 10.2 for controls with favorable local histology.

# **Analyses: sampling weights**

### Effect of unfavorable histology

	<u> </u>	
	log OR	SE
Full data	1.81	0.11
Subcohort		
(ignoring sampling)	-0.02	0.12
two-phase	1.78	0.155
single-phase	1.78	0.159

### Can use any analysis, eg RR rather than OR

	log RR	SE
Full data	1.39	0.07
Subcohort		
(ignoring sampling)	-0.01	0.06
two-phase	1.36	0.101
single-phase	1.36	0.107

Declaring a two-phase design

- Data set has records for all phase-one people, subset variable indicates membership in second phase
- Two id, two strata.
- Second-phase weights and fpc are worked out by R

We can compare to the conservative approximation: single-phase sampling with replacement

Almost no advantage of two-phase analysis in this example; but wait until this afternoon and calibration of weights.

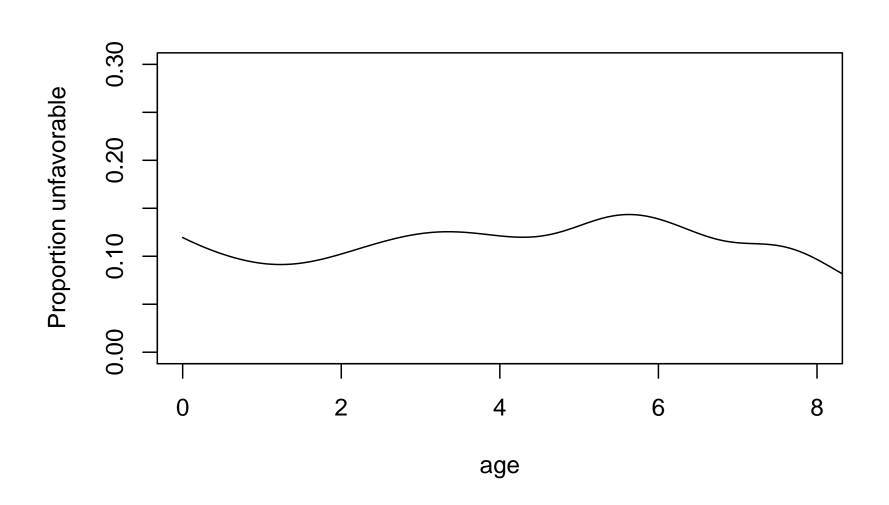
```
summary(svyglm(rel~factor(stage)*factor(histol),design=dccs2,
                        family=quasibinomial()))
                            Estimate Std. Error t value Pr(>|t|)
(Intercept)
                               -2.6946
                                           0.1246 -21.623 < 2e-16 ***
factor(stage)2
                                0.7733
                                           0.1982
                                                    3.901 0.000102 ***
factor(stage)3
                                0.7400
                                           0.2048
                                                  3.614 0.000315 ***
factor(stage)4
                                           0.2572 4.401 1.18e-05 ***
                                1.1322
factor(histol)2
                                1.1651
                                           0.3196 3.646 0.000279 ***
factor(stage)2:factor(histol)2
                                0.3642
                                           0.4462 0.816 0.414511
factor(stage)3:factor(histol)2
                                1.0230
                                           0.3968 2.578 0.010056 *
factor(stage)4:factor(histol)2
                                1.7444
                                           0.4973
                                                    3.508 0.000470 ***
  summary(svyglm(rel~factor(stage)*factor(histol),design=dcons,
                        family=quasibinomial()))
                              Estimate Std. Error t value Pr(>|t|)
(Intercept)
                               -2.6946
                                           0.1355 - 19.886 < 2e - 16 ***
factor(stage)2
                                0.7733
                                           0.1982
                                                    3.902 0.000101 ***
factor(stage)3
                                0.7400
                                           0.2047 3.615 0.000314 ***
factor(stage)4
                                1.1322
                                           0.2571 4.403 1.17e-05 ***
factor(histol)2
                                1.1651
                                           0.3208 3.632 0.000294 ***
factor(stage)2:factor(histol)2
                                0.3642
                                           0.4461 0.816 0.414408
factor(stage)3:factor(histol)2
                                1.0230
                                           0.3974 2.574 0.010169 *
factor(stage)4:factor(histol)2
                                1.7444
                                           0.4977
                                                    3.505 0.000474 ***
```

We are not restricted to logistic regression: eg, log link gives log relative risks rather than log odds ratios. SInce relapse is not rare for unfavorable histology these are noticeably different.

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                                         0.1167 - 23.644 < 2e-16 ***
                              -2.7600
factor(stage)2
                               0.7020
                                         0.1790 3.922 9.30e-05 ***
factor(stage)3
                               0.6730
                                         0.1849 3.640 0.000285 ***
factor(stage)4
                               1.0072
                                         0.2207 4.564 5.58e-06 ***
factor(histol)2
                               1.0344
                                         0.2690 3.845 0.000127 ***
factor(stage)2:factor(histol)2
                               0.1153
                                         0.3405
                                                 0.339 0.734920
factor(stage)3:factor(histol)2
                               0.4695
                                         0.3118 1.505 0.132495
factor(stage)4:factor(histol)2
                               0.4872
                                         0.3316
                                                 1.469 0.142020
```

Other exploratory and descriptive analyses are also possible: how does histology vary with age at diagnosis?

[Really needs more detailed classification to be useful analysis]



# Case-cohort design

Cox model estimation involves solving

$$\sum_{\text{event}} Z_{\text{event}} - E(\beta, t) = 0$$

where  $Z_{\text{event}}$  is the covariate for the person having an event and  $E(\beta,t)$  is the expected covariate based on a weighted average over everyone at risk at that time.

## Case-cohort design

- Prentice (1986) suggested estimating  $E(\beta,t)$  from the current case plus a subcohort selected at random at baseline, saving money and computation time.
- Self & Prentice suggested dropping current case from  $E(\beta,t)$  for mathematical simplicity.
- Barlow used the subcohort and all cases, with the conservative single-phase sampling standard errors
- Lin & Ying used the same estimates as Barlow, but the correct standard errors.
- Borgan et al (2000) allowed stratified sampling and timedependent sampling weights.

## **Advantages**

Case—cohort and nested case—control studies have similar power, but in a case—cohort design

- The same subcohort can be used for more than one outcome and for different lengths of followup
- Subcohort defined at baseline allows exposure measurement spread over whole study (for some measurements)
- Information from whole cohort can easily be incorporated when selecting subcohort (eg surrogate exposure) or with post-stratification at time of analysis

Design has been used in ARIC, CHS for measuring genotypes or biomarkers.

### Historical artifact

Mathematical techniques in 1980s,1990s, required viewing the Cox model estimating equations

$$\sum_{\text{event}} Z_{\text{event}} - E(\beta) = 0$$

as a process evolving over time.

A member of the subcohort who was in the future going to be a case still had to be treated the same as someone who was not going to become a case, because that information was not predictable. Weight might be 10 until the event and then 1 at the event.

Survey techniques allow a simpler view *sub specie aeternitatis*; no hang-ups about predictable weight processes: weight of 1 for any future case.

# Computation

- Prentice, Self & Prentice methods are not just weighted Cox regression, so need some trickery. Mayo Clinic Biostatistics Technical Report #62 describes the trickery and gives S-PLUS and SAS code fragments. cch() in R survival package is based on these methods.
- Barlow's method, using the conservative single-phase approximateion, is just weighted Cox regression and works in any software with survey or robust standard errors.
- Lin & Ying's method is two-phase weighted Cox regression, needs software for two-phase analysis.

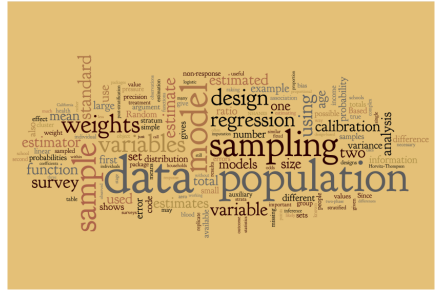
```
Estimators based on survey-weighted Cox models (Lin & Ying)
dcchs<-twophase(id=list(~seqno,~seqno), strata=list(NULL,~rel),</pre>
              subset=~I(in.subcohort | rel), data=nwtco)
svycoxph(Surv(edrel,rel)~factor(stage)+factor(histol)+I(age/12),
              design=dcchs)
Traditional estimators based on the Therneau & Li trick.
fit.ccSP <- cch(Surv(edrel, rel) ~ stage + histol + age,
               data =ccoh.data, subcoh = ~subcohort, id=~seqno,
               cohort.size=4028, method="SelfPren")
stratsizes<-table(nwtco$instit)
fit.BI<- cch(Surv(edrel, rel) ~ stage + histol + age,
            data =ccoh.data, subcoh = ~subcohort,
            id=~seqno, stratum=~instit, cohort.size=stratsizes,
            method="I.Borgan")
```

If you liked today's talk and want to read more

or

If you didn't like today's talk and want a book instead

Complex Surveys: a guide to analysis using R coming soon from Wiley



http://faculty.washington.edu/tlumley/svybook/