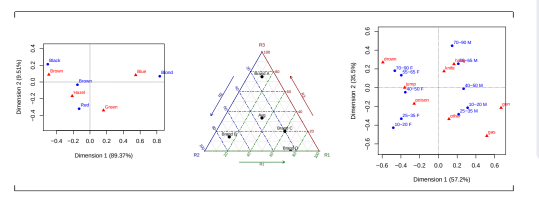
# Correspondence analysis

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Basic ideas

# Correspondence analysis: Basic ideas

#### **Correspondence analysis (CA)**

Analog of PCA for frequency data:

- account for maximum % of  $\chi^2$  in few (2-3) dimensions
- finds scores for row  $(x_{im})$  and column  $(y_{jm})$  categories on these dimensions
- uses Singular Value Decomposition of residuals from independence,  $d_{ij}=(n_{ij}-\widehat{m}_{ij})/\sqrt{\widehat{m}_{ij}}$

$$d_{ij} = \sqrt{n} \sum_{m=1}^{M} \lambda_m x_{im} y_{jm} \quad \leftrightarrow \quad \mathbf{D} = \mathbf{X} \mathbf{\Lambda} \mathbf{Y}^{\mathsf{T}}$$

- *optimal scaling*: each pair of scores for rows  $(x_{im})$  and columns  $(y_{jm})$  have highest possible correlation  $(= \lambda_m)$ .
- plots of the row  $(x_{im})$  and column  $(y_{im})$  scores show associations

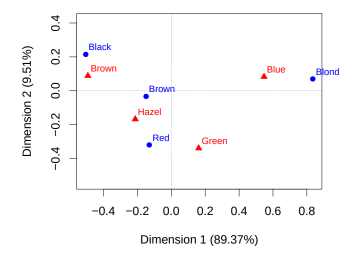
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# Example: Hair color, eye color data

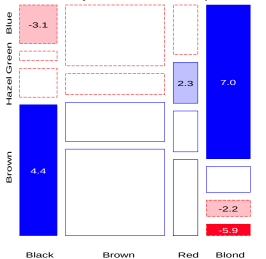
```
library(ca)
(haireye.ca <- ca(haireye))
    Principal inertias (eigenvalues):
             0.208773 0.022227 0.002598
  Percentage 89.37%
                      9.52%
    Rows:
                    Brown
                   0.4831
           0.1824
  ChiDist 0.5512
                   0.1595
                          0.3548 0.838
          0.0554
                  0.0123 0.0151 0.151
       1 -1.1043 -0.3245 -0.2835 1.828
          1.4409 -0.2191 -2.1440 0.467
    Columns:
            Brown Blue
                          Hazel
           0.3716 0.363
   ChiDist 0.5005 0.554
  Inertia 0.0931 0.111 0.0131
  Dim. 1 -1.0771 1.198 -0.4653 0.3540
          0.5924 0.556 -1.1228 -2.2741
```

Hair color, Eye color data:



- Rough interpretation: row/column points "near" each other are positively associated
- Dim 1: 89.4% of  $\chi^2$  (dark  $\leftrightarrow$  light)
- Dim 2: 9.5% of  $\chi^2$  (Red/Green vs. others)

Hair color, Eye color data: Compare with mosaic display



Row and column profiles

- The main dark-light dimension is reflected in the opposite-corner pattern of residuals
- The 2nd dimension is reflected in deviations from this pattern (e.g., Red hair-Green eyes)
- CA is "accounting for" residuals (deviations) from independence

### Row and column profiles

- For a two-way table, row profiles and column profiles give the relative proportions of the column/row categories.
- An association is present to the extent that the row/col profiles differ
- Profiles add to 1.0 (100%), and can be visualized in profile space

#### **Example: Toothpaste purchases by region**

120 people in three regions where asked which of four brands of toothpaste, A-D, they had most recently purchased. Is there a difference among regions?

```
toothpaste
            Region
## Brand
             R1 R2 R3
     Brand A 5
     Brand B 5 25
     Brand C 15
     Brand D 15
```

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Plotting profiles

Row profiles pertain to the differences among brand preference

Column profiles pertain to the differences among regions

```
Brand A 12.5 12.5 75.0 100
                                              12.5
  Brand B 14.3 71.4 14.3 100
                                    ## Brand B
## Brand C 60.0 20.0 20.0 100
                                               37.5
                                   ## Brand C
                                                     12.5
## Brand D 75.0 25.0 0.0 100
                                    ## Brand D
                                              37.5 12.5
                                                            0.0
          33.3 33.3 33.3 100
                                              100.0 100.0 100.0 100.0
```

There is clearly an association, meaning that the row (column) profiles differ

```
chisq.test(toothpaste)
   Pearson's Chi-squared test
## data: toothpaste
\#\# X-squared = 79.6, df = 6, p-value = 4.3e-15
```

In this simple example we can plot the row profiles as points in 3D space, with axes corresponding to regions, R1, R2, R3

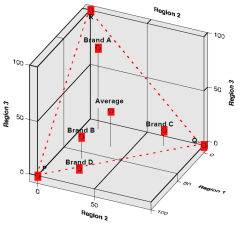
Profiles

• Each brand is positioned in this space according to its proportions for the regions

 Because proportions sum to 100%, all points lie in the dashed plane PQR

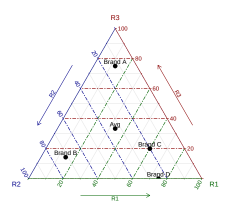
 The Average profile is at the (weighted) centroid

 If no association, all brands would appear at the centroid



# Plotting profiles

Analogous 2D plot is a trilinear plot that automatically scales the R1–R3 values so they sum to 100%



- The Avg profile has coordinates of 33.3% for each region
- Brand preferences by region can be seen by their positions wrt the R1-R3 axes
- This suggests that differences among brands can be measured by their (squared) distances from the centroid, weighted by their row margins (mass)
- Physical analogy suggests the term inertia for this weighted variation

### **CA** solution

- The CA solution has at most min(r-1, c-1) dimensions
- A 2D solution here is exact, i.e., accounts for 100% of Pearson  $X^2$

```
library(ca)
tp.ca <- ca(toothpaste)
summary(tp.ca, rows=FALSE, columns=FALSE)
##
##
  Principal inertias (eigenvalues):
##
    dim
           value
                           cum%
                                   scree plot
##
           0.410259
                      61.8
                            61.8
##
           0.253134
                      38.2 100.0
##
    Total: 0.663393 100.0
```

#### Pearson $X^2$ :

```
sum(tp.ca$sv^2) * 120
## [1] 79.607
```

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Basic ideas

Profiles

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### **CA** solution

```
res <- plot(tp.ca)
polygon(res$cols, border="red", lwd=2)
```

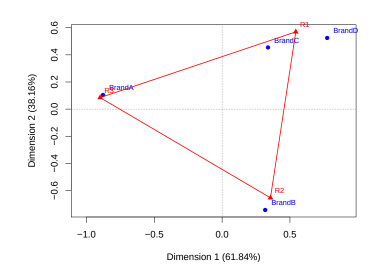
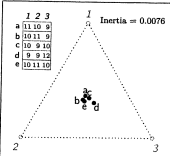
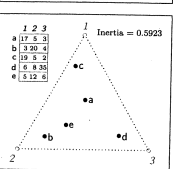
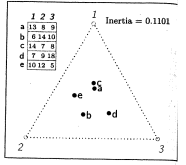


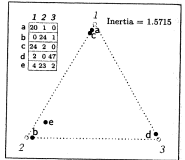
Exhibit 4.2: A series of data tables with increasing total inertia. The higher the total inertia, the greater is the association between the rows and columns, displayed by the higher dispersion of the profile points in the profile space. The values in these tables have been chosen specifically so that the column sums are all equal. so the weights in the  $\chi^2$ -distance formulation are the same, and hence distances we observe in these maps are

true  $\chi^2$ -distances.







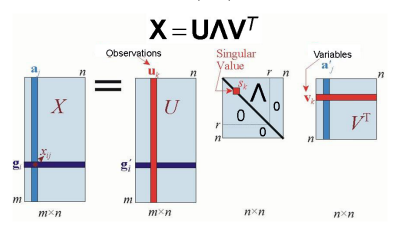


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# Singular value decomposition

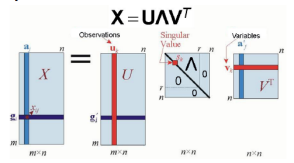
The singular value decomposition (SVD) is a basic technique for factoring a matrix and matrix approximation.

For an  $m \times n$  matrix  $\boldsymbol{X}$  of rank  $r \leq \min(m, n)$  the SVD of  $\boldsymbol{X}$  is:



### Properties of the SVD

- V: columns are the eigenvectors of X<sup>T</sup>X and form an orthonormal basis (V<sup>T</sup>V = I) for the variables
- Λ: diagonal, r singular values are the square roots of the eigenvalues of both XX<sup>T</sup> and X<sup>T</sup>X
- U: columns are the eigenvectors of XX<sup>T</sup> and form an orthonormal basis for the observation profiles, so that U<sup>T</sup>U = I



CA coordinates

SVD: Matrix approximation

• Let **X** be an  $m \times n$  matrix such that rank(X) = r

Basic ideas

• If  $\lambda_1 \geq \lambda_2 \geq ... \geq \lambda_r$  are the singular values of **X**, then  $\hat{\mathbf{X}}$ , the rank q approximation of **X** that minimizes  $||\mathbf{X} - \hat{\mathbf{X}}||$ , is

$$\hat{\mathbf{X}}_{m \times n} = \sum_{i=1}^{q} \lambda_i \begin{pmatrix} u_{i1} \\ \vdots \\ u_{im} \end{pmatrix} \begin{pmatrix} \mathbf{v}_{i1} & \cdots & \mathbf{v}_{in} \end{pmatrix} = \lambda_1 u_1 \mathbf{v}_1^T + \cdots + \lambda_q u_q \mathbf{v}_q^T$$
row scores

a sum of q rank=1 (outer) products. The variance in **X** accounted for each term is  ${\lambda_1}^2$ 

# CA notation and terminology

#### Notation:

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- Contingency table:  $N = \{n_{ii}\}$
- Correspondence matrix (cell probabilities):  $\mathbf{P} = \{p_{ii}\} = \mathbf{N}/n$
- Row/column masses (marginal probabilities):  $r = \sum_i p_{ij}$  and  $c = \sum_i p_{ij}$
- Diagonal weight matrices:  $\mathbf{D}_r = \operatorname{diag}(\mathbf{r})$  and  $\mathbf{D}_c = \operatorname{diag}(\mathbf{c})$

The SVD is then applied to the correspondence matrix of cell probabilities as:

$$P = AD_{\lambda}B^{\mathsf{T}}$$

#### where

- Singular values:  $\mathbf{D}_{\lambda} = \operatorname{diag}(\lambda)$  is the diagonal matrix of singular values  $\lambda_1 \geq \lambda_2 \geq \cdots \geq \lambda_M$
- Row scores:  $\mathbf{A}_{I\times M}$ , normalized so that  $\mathbf{A}\mathbf{D}_{I}^{-1}\mathbf{A}^{\mathsf{T}}=\mathbf{I}$
- Column scores:  $\mathbf{B}_{J\times M}$ , normalized so that  $\mathbf{B}\mathbf{D}_c^{-1}\mathbf{B}^{\mathsf{T}}=\mathbf{I}$

# Principal and standard coordinates

Two types of coordinates are commonly used in CA, based on re-scalings of **A** and **B**.

#### **Principal coordinates**

Coordinates of the row (F) and column (G) profiles wrt their own principal axes

$$F = D_r^{-1}AD_{\lambda}$$
 scaled so that  $F^TD_rF = D_{\lambda}$   
 $G = D_c^{-1}BD_{\lambda}$  scaled so that  $G^TD_cG = D_{\lambda}$ 

- Defined so that the inertia along each axis is the corresponding singular value,  $\lambda_i$ ,
- i.e., weighted average of squared principal coordinates =  $\lambda_i$  on dim. i
- The joint plot in principal coordinates, F and G, is called the symmetric map because both row and column profiles are overlaid in the same coordinate system.

CA coordinates

### Principal and standard coordinates

#### Standard coordinates

The standard coordinates  $(\Phi,\Gamma)$  are a rescaling of the principal coordinates to unit inertia along each axis,

$$\Phi = \mathbf{D}_r^{-1}\mathbf{A}$$
 scaled so that  $\Phi^T \mathbf{D}_r \Phi = \mathbf{I}$   
 $\Gamma = \mathbf{D}_c^{-1}\mathbf{B}$  scaled so that  $\Gamma^T \mathbf{D}_c \Gamma = \mathbf{I}$ 

- The weighted average of squared standard coordinates = 1 on each dimension
- An asymmetric map shows one set of points (say, the rows) in principal coordinates and the other set in standard coordinates.

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Basic ideas

### Geometric and statistical properties

nested solutions: CA solutions are *nested*, meaning that the first two dimensions of a 3D solution will be identical to the 2D solution (similar to PCA)

Basic ideas

centroids at the origin: In both principal coordinates and standard coordinates the points representing the row and column profiles have their centroids (weighted averages) at the origin. The origin represents the (weighted) average row profile and column profile.

chi-square distances: In principal coordinates, the row coordinates are equal to the row profiles  $\mathbf{D}_r^{-1}\mathbf{P}$ , rescaled inversely by the square-root of the column masses,  $\mathbf{D}_c^{-1/2}$ . Distances between two row profiles,  $\mathbf{R}_i$  and  $\mathbf{R}_{i'}$  are  $\chi^2$  distances, where the squared difference  $[\mathbf{R}_{ij} - \mathbf{R}_{i'j}]^2$  is inversely weighted by the column frequency, to account for the different relative frequency of the column categories.

### The ca package in R

• ca () calculates CA solutions, returning a "ca" object

ca package

The result contains the standard row coordinates (rowcoord: Φ) and column coordinates (colcoord: Γ)

```
haireye.ca$rowcoord

## Dim1 Dim2 Dim3

## Black -1.10428 1.44092 -1.08895

## Brown -0.32446 -0.21911 0.95742

## Red -0.28347 -2.14401 -1.63122

## Blond 1.82823 0.46671 -0.31809
```

c ideas ca package Basic ideas ca package

# The ca package in R

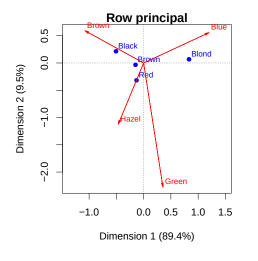
The plot () method provides a wide variety of scalings (map=), with different interpretive properties. Some of these are:

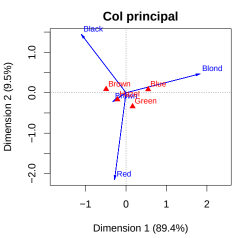
- "symmetric" both rows and columns in pricipal coordinates (default)
- "rowprincipal" or "colprincipal" asymmetric maps, with either rows in principal coordinates and columns in standard coordinates, or vice versa
- "symbiplot" scales both rows and columns to have variances equal to the singular value

The mcja() function is used for multiple correspondence analysis (MCA) and has analogous print(), summary() and plot() methods.

Asymmetric row/col principal plots are biplots — can interpret projection of points on vectors

```
plot (haireye.ca, map="rowprincipal", arrows=c(FALSE, TRUE))
plot (haireye.ca, map="colprincipal", arrows=c(TRUE, FALSE))
```





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CA properties

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### Optimal category scores

- CA has a close relation to canonical correlation analysis, applied to dummy variables representing the categories.
- The singular values,  $\lambda_i$ , are the correlations between the category scores
  - Assign Dim 1 scores, X1 and Y1 to the row/column categories:  $\implies$  max. possible correlation,  $\lambda_1$
  - Assign Dim 2 scores, X2 and Y2 to the row/column categories:  $\implies$  max. possible correlation,  $\lambda_2$ , but uncorrelated with X1, Y1
  - Thus all association between the row/col categories is captured by the scores
- This optimal scaling interpretation can be used to quantify categorical variables

### Optimal category scores

#### Singular values = canonical correlations

```
haireye.ca <- ca(haireye)
round(haireye.ca$sv, 4)
## [1] 0.4569 0.1491 0.0510
```

#### Extract the row and column coordinates to a data frame

```
RC <- haireye.ca$rowcoord # row coordinates
CC <- haireye.ca$colcoord # col coordinates
HE.df <- as.data.frame(haireye)

Y1 <- RC[match(HE.df$Hair, haireye.ca$rownames),1]
X1 <- CC[match(HE.df$Eye, haireye.ca$colnames),1]
Y2 <- RC[match(HE.df$Hair, haireye.ca$rownames),2]
X2 <- CC[match(HE.df$Eye, haireye.ca$colnames),2]
```

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# Optimal category scores

```
HE.df <- cbind(HE.df, X1, Y1, X2, Y2)
print (HE.df, digits=3)
      Hair
             Eye Freq
                           Х1
                                  Υ1
                                         Х2
                    68 -1.077 -1.104
                                      0.592
                                             1.441
     Black Brown
     Brown Brown
                  119 -1.077 -0.324
                    26 -1.077 -0.283
                                      0.592 - 2.144
        Red Brown
     Blond Brown
                    7 -1.077 1.828
                                      0.592
            Blue
                    20 1.198 -1.104
     Brown
            Blue
                    84 1.198 -0.324
                                      0.556 - 0.219
            Blue
                   17
                       1.198 -0.283
                                      0.556 - 2.144
        Red
     Blond
            Blue
                    94 1.198 1.828 0.556 0.467
```

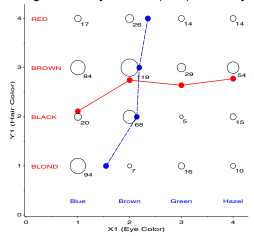
#### Calculate correlations—all zero except $r(X1,Y1)=\lambda_1$ and $r(X2,Y2)=\lambda_2$

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CA properties

### Simultaneous linear regressions

Assign arbitrary scores (1–4) X1 to eye color and Y1 to hair color

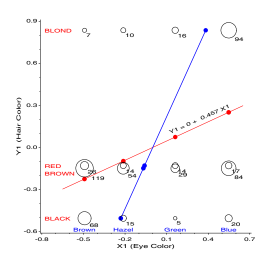


- Lines connecting the weighted (conditional) means of Y1 | X1 and X1 | Y1 are not-linear
- The scatterplot uses bubble symbols showing frequency in each cell
- Is it possible to assign row and column scores so that both regressions are linear?

CA examples

# Simultaneous linear regressions

#### Yes, use CA scores on the first dimension



- The regression of Y1 on X1 is linear, with slope  $\lambda_1$
- The regression of X1 on Y1 is linear, with slope  $1/\lambda_1$
- λ<sub>1</sub> is the (canonical) correlation between X1 and Y1
- The angle between the two lines would be 0 if perfect correlation
- The conditional means (dots) are the principal coordinates

# Example: Mental impairment and parents' SES

Data on mental health status (mental) of 1660 young NYC residents by parents' SES (ses), a  $6 \times 4$  table.

- Both mental and ses are ordered factors
- Convert from frequency data frame to table using xtabs ()

```
data("Mental", package="vcdExtra")
str(Mental)

## 'data.frame': 24 obs. of 3 variables:
## $ ses : Ord.factor w/ 6 levels "1"<"2"<"3"<"4"<...: 1 1 1 1 2 2 2 2 2
## $ mental: Ord.factor w/ 4 levels "Well"<"Mild"<...: 1 2 3 4 1 2 3 4 1
## $ Freq : int 64 94 58 46 57 94 54 40 57 105 ...
mental.tab <- xtabs(Freq ~ ses + mental, data=Mental)</pre>
```

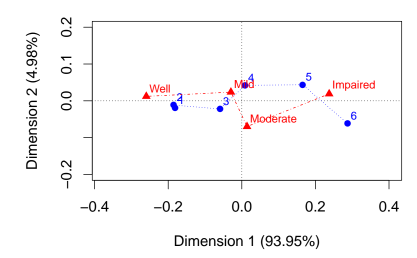
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examples CA examples

# Example: Mental impairment and parents' SES

- The exact CA solution has min(r-1, c-1) = 3 dimensions
- The total Pearson  $X^2$  is  $n\Sigma\lambda_i^2 = 1660 \times 0.0277 = 45.98$  with 15 df
- Of this, 93.9% is accounted for by the first dimension

```
res <- plot (mental.ca)
lines (res$rows, col="blue", lty=3)
lines (res$cols, col="red", lty=4)</pre>
```



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Multi-way tables

CA examples

### Looking ahead

- CA is largely an exploratory method row/column scores are not parameters of a statistical model; no confidence intervals
- Only rough tests for the number of CA dimensions
- Can't test an hypothesis that the row/column scores are have some particular spacing (e.g., are mental and ses equally spaced?)
- These kinds of questions can be answered with specialized loglinear models
- Nevertheless, plot (ca (table)) gives an excellent quick view of associations

# Multi-way tables

Correspondence analysis can be extended to *n*-way tables in several ways:

#### Stacking approach

- *n*-way table flattened to a 2-way table, combining several variables "interactively"
- Each way of stacking corresponds to a loglinear model
- ullet Ordinary CA of the flattened table o visualization of that model
- Associations among stacked variables are not visualized

#### Multiple correspondence analysis (MCA)

- Extends CA to n-way tables
- Analyzes all pairwise bivariate associations
- Can plot all factors in a single plot
- An extension, joint correspondence analysis, gives a better account of inertia for each dimension

3.

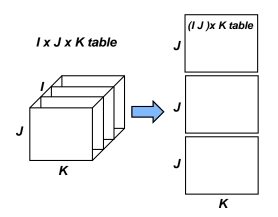
Multi-way tables Stacking Multi-way tables

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# Multi-way tables: Stacking

#### Stacking approach:

• three-way table, of size  $I \times J \times K$  can be sliced and stacked as a two-way table in different ways



- The variables combined are treated "interactively"
- Each way of stacking corresponds to a loglinear model
  - (I × J) × K → [AB][C]
  - $I \times (J \times K) \rightarrow [A][BC]$
  - $J \times (I \times K) \rightarrow [B][AC]$
- Only the associations in separate [] terms are analyzed and displayed
- The stacked table is analyzed with ordinary CA of the two-way stacked table

### Interactive coding in R

• Data in table (array) form: Use as.matrix(structable())

```
mat1 <- as.matrix(structable(A + B ~ C, data=mytable)) # [A B][C]</pre>
mat2 <- as.matrix(structable(A + C ~ B + D, data=mytable)) # [A C][B I</pre>
ca (mat2)
```

• Data in frequency data frame form: Use paste() or interaction(), followed by xtabs ()

```
mydf$AB <- interaction(mydf$A, mydf$B, sep='.')</pre>
                                                       # levels: A.B
mydf$AB <- paste(mydf$A, mydf$B, sep=':')</pre>
                                                       # levels: A:B
mytab <- xtabs(Freq ~ AB + C, data=mydf)</pre>
                                                       # [A B] [C}
```

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Multi-way tables Stacking

Multi-way tables Stacking

# Example: Suicide rates in Germany

- Suicide in vcd gives a  $2 \times 5 \times 8$  table of sex by age.group by method of suicide for 53,182 suicides in Germany, in a frequency data frame
- With the data in this form, you can use paste () to join age.group and sex together to form a new variable age sex consisting of their combinations.

```
data("Suicide", package="vcd")
# interactive coding of sex and age.group
Suicide <- within (Suicide, {
        age_sex <- paste(age.group, toupper(substr(sex,1,1)))</pre>
```

# Example: Suicide rates in Germany

```
suicide.tab <- xtabs(Freq ~ age sex + method2, data=Suicide)</pre>
suicide.tab
            method2
## age_sex
             poison
                     gas hang drown
                                     gun knife jump other
    10-20 F
                                      25
                921
                      40 212
                                 30
                                             11
                                                 131
                                                       100
    10-20 M
               1160
                     335 1524
                                 67
                                     512
                                             47
                                                 189
                                                       464
    25-35 F
               1672 113 575
                                      64
                                                 276
                                                       263
                                139
                                             41
    25-35 M
               2823
                     883 2751
                                213
                                     852
                                            139
                                                 366
                                                       775
     40-50 F
                      91 1481
                                354
                                      52
                                             80
                                                 327
                                                       305
               2224
     40-50 M
                     625 3936
                                247
                                     875
                                           183
                                                 244
                                                       534
               2465
     55-65 F
               2283
                      45 2014
                                679
                                            103
                                                388
                                                       296
                                     477
##
     55-65 M
               1531
                     201 3581
                                207
                                            154 273
                                                       294
     70-90 F
               1548
                      29 1355
                                501
                                      3
                                             74
                                                 383
                                                       106
     70-90 M
                938
                      45 2948
                                212 229
                                            105 268
                                                       147
```

- The CA analysis will be that of the loglinear model [Age Sex] [Method]
- It will show associations between the age—sex combinations and method of suicide
- Associations between age and sex will not be shown in this analysis

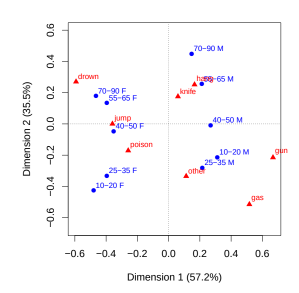
Multi-way tables Stacking Multi-way tables Stacking

# Example: Suicide rates in Germany

```
suicide.ca <- ca(suicide.tab)</pre>
summary(suicide.ca)
  Principal inertias (eigenvalues):
    dim
           value
                      응
                           cum%
                                  scree plot
           0.096151 57.2
                           57.2
           0.059692 35.5
                           92.6
           0.008183
                     4.9
                           97.5
           0.002158
                      1.3
                           98.8
           0.001399
                      0.8 99.6
           0.000557
                      0.3 100.0
           6.7e-050
                    0.0 100.0
   Total: 0.168207 100.0
```

It can be seen that 92.6% of the Pearson  $X^2$  for this model is accounted for in the first two dimensions.

```
plot(suicide.ca)
```

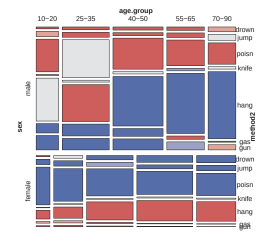


- Dim 1: Sex
- Dim 2: Age
- Interpret method use by age-sex combinations, e.g., young males: gas, gun; young

females: poison

#### Multi-way tables Compare with a mosaic plot also fitting the model [Age Sex][Suicide]:

```
suicide.tab3 <- xtabs(Freq ~ sex + age.group + method2, data=Suicide)</pre>
mosaic(suicide.tab3, shade=TRUE, legend=FALSE,
       expected=~age.group*sex + method2,
       labeling_args=list(abbreviate_labs=c(FALSE, FALSE, 5)),
                           rot_labels = c(0, 0, 0, 90))
```



### Marginal tables and supplementary variables

• An *n*-way table is collapsed to a marginal table by ignoring factors

Multi-way tables

Omitted variables can be included by treating them as supplementary

Marginal tables

These are projected into the space of the marginal CA

Age by method, ignoring sex:

```
suicide.tab2 <- xtabs(Freq ~ age.group + method2, data=Suicide)</pre>
suicide.tab2
            method2
                     gas hang drown
                                      gun knife jump other
## age.group poison
       10-20
               2081
                     375 1736
                                 97
                                      537
                                             58
                                                 320
                                                        564
##
                                                 642 1038
       25-35
               4495
                     996 3326
                                 352
                                      916
                                            180
                                            263 571
       40 - 50
               4689 716 5417
                                 601
                                      92.7
                                                        839
       55-65
               3814
                     246 5595
                                 886
                                      506
                                            257
                                                 661
       70-90
               2486
                     74 4303
                                 713
                                      232
                                            179
                                                 651
                                                        253
```

#### Relation of sex and method:

```
(suicide.sup <- xtabs(Freq ~ sex + method2, data=Suicide))
           method2
## sex
            poison
                     gas hang drown
                                       qun knife
                                                   jump other
              8917
                    2089 14740
                                 946
                                      2945
                                              628
                                                  1340
                                                         2214
    male
                    318 5637 1703
    female
              8648
                                       173
                                              309
                                                  1505
suicide.tab2s <- rbind(suicide.tab2, suicide.sup)</pre>
```

# Marginal tables and supplementary variables

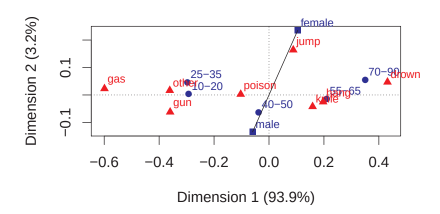
The rows for sex by method are treated as supplementary rows:

```
suicide.ca2s <- ca(suicide.tab2s, suprow=6:7)</pre>
summary(suicide.ca2s)
## Principal inertias (eigenvalues):
##
   dim
           value
                      % cum%
                                 scree plot
   1
           0.060429 93.9 93.9
           0.002090
           0.001479
                      2.3 99.4
           0.000356 0.6 100.0
   Total: 0.064354 100.0
##
```

- the relation of age and method is now essentially 1 dimensional
- the inertia of Dim 1 (0.604) is nearly the same as that of Dim 2 (0.596) in the stacked table

- Plot the 2D CA solution for the [Age] [Method] marginal table.
- Add category points for Sex (supplementary points)

res <- plot(suicide.ca2s, pch=c(16, 15, 17, 24)) lines(res\$rows[6:7,])



Ignoring Sex has collapsed Dim 1 (Sex) of the [Age Sex][Method] analysis. Supp. points show associations of Method with Sex.

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Multiple correspondence analysis

Multiple correspondence analysis

# Multiple correspondence analysis (MCA)

- Extends CA to n-way tables
- Useful when simpler stacking approach doesn't work well, e.g., 10 categorical attitude items
- Analyzes all pairwise bivariate associations. Analogous to:
  - Correlation matrix (numbers)
  - Scatterplot matrix (graphs)
  - All pairwise  $\chi^2$  tests (numbers)
  - Mosaic matrix (graphs)
- Provides an optimal scaling of the category scores for each variable
- Can plot all factors in a single plot
- An extension, joint correspondence analysis, gives a better account of inertia for each dimension

### The indicator matrix and the Burt matrix

Two ways to think about MCA:

#### Indicator matrix (dummy variables)

- A given categorical variable, q, can be represented by an indicator matrix  $\mathbf{Z}(n \times J_q)$  of dummy variables,  $z_{ii} = 1$  if case i is in category j
- Let  $Z_1, Z_2, \dots, Z_Q$  be the indicator matrices for Q variables
- MCA is then a simple CA applied to the partitioned matrix  $\mathbf{Z} = [\mathbf{Z}_1, \mathbf{Z}_2, \dots, \mathbf{Z}_Q]$

#### **Burt matrix**

• The Bert matrix is the product of the indicator matrix **Z** and its transpose

$$\boldsymbol{B} = \boldsymbol{Z}^{\mathsf{T}} \boldsymbol{Z}$$

 MCA can be defined using the SVD of B, giving category scores for all variables accounting for the largest proportion of all bivariate associations.

Multiple correspondence analysis Bivariate MCA Multiple correspondence analysis Bivariate MCA

# Bivariate MCA: Hair Eye color

- For the hair-eye color data, the indicator matrix Z has n=592 rows (observations) and 4 + 4 = 8 columns (categories).
- Shown below in frequency form, using h1-h4 for hair color and e1-e4 for eye color
- E.g., first row reflects the 68 observations with black hair and brown eyes

```
Eye Freq h1 h2 h3 h4 e1 e2 e3 e4
     Black Brown
                  68
     Brown Brown
                119
## 3
                 26
                     0
                        0
                          1
       Red Brown
    Blond Brown
                        0
                          0
    Black Blue
                 20
                     1
     Brown Blue
                 84
                        1
       Red Blue
                17
                     0 0 1 0 0
    Blond Blue
                 94 0
                        0 0 1
```

#### Expand this to case form for **Z** (592 x 8)

```
Z <- expand.dft(haireye.df)[,-(1:2)]
vnames <- c(levels(haireye.df$Hair), levels(haireye.df$Eye))
colnames(Z) <- vnames
dim(Z)
## [1] 592 8</pre>
```

If the indicator matrix is partitioned as  $Z = [Z_1, Z_2]$ , corresponding to the hair, eye categories, then the contingency table is given by  $N = Z_1^T Z_2$ .

```
(N \leftarrow t(as.matrix(Z[,1:4])) %*% as.matrix(Z[,5:8]))
          Brown Blue Hazel Green
## Black
             68
                   20
                         15
                         54
                                29
## Brown
            119
             26
## Red
                  17
                         14
                                14
## Blond
                   94
                         10
                                16
```

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Multiple correspondence analysis

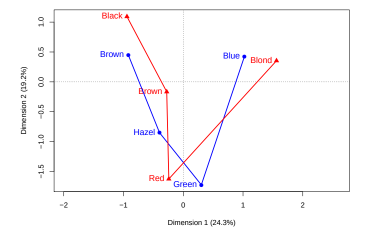
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Multiple correspondence analysis

sis Bivariate MCA

- We can then use ordinary CA on the indicator matrix, Z
- Except for scaling, this is the same as the CA of N
- The inertia contributions differ, and this is handled better by MCA



### The Burt matrix

For two categorical variables, the Burt matrix is

$$m{B} = m{Z}^{\mathsf{T}} m{Z} = \left[ egin{array}{ccc} m{N}_1 & m{N} \ m{N}^{\mathsf{T}} & m{N}_2 \end{array} 
ight] \;\; .$$

- $N_1$  and  $N_2$  are diagonal matrices containing the marginal frequencies of the two variables
- The contingency table, **N** appears in the off-diagonal block

A similar analysis to that of the indicator matrix Z is produced by:

```
Burt <- t(as.matrix(Z)) %*% as.matrix(Z)
rownames(Burt) <- colnames(Burt) <- vnames
Burt.ca <- ca(Burt)
plot(Burt.ca)</pre>
```

- Standard coords are the same
- Singular values of **B** are the squares of those of **Z**

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### Multivariate MCA

For Q categorical variables, the Burt matrix is

$$m{B} = m{Z}^{\mathsf{T}} m{Z} = \left[ egin{array}{cccc} m{N}_1 & m{N}_{[12]} & \cdots & m{N}_{[1Q]} \ m{N}_{[21]} & m{N}_2 & \cdots & m{N}_{[2Q]} \ dots & dots & \ddots & dots \ m{N}_{[Q1]} & m{N}_{[Q2]} & \cdots & m{N}_Q \end{array} 
ight] \; .$$

- $\bullet$  The diagonal blocks  $N_i$  contain the one-way marginal frequencies
- The off-diagonal blocks  $N_{[ij]}$  contain the bivariate contingency tables for each pair (i, j) of variables.
- Classical MCA can be defined as a SVD of the matrix B
- It produces scores for the categories of *all* variables accounting for the greatest proportion of the bivariate associations in off-diagonal blocks in a small number of dimensions.

### MCA properties

- The inertia contributed by a given variable increases with the number of response categories: inertia ( $\mathbf{Z}_a$ ) =  $J_a$  1
- The centroid of the categories for each variable is at the origin of the display.
- For a given variable, the inertia contributed by a given category increases as the marginal frequency in that category *decreases*. Low frequency points therefore appear further from the origin.
- The category points for a binary variable lie on a line through the origin.

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Multiple correspondence analysis

MCA example

Multiple correspondence analysis

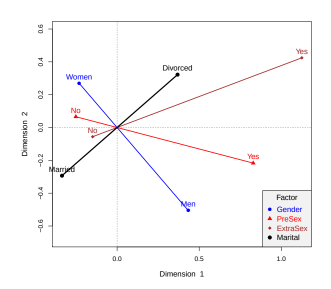
MCA ov

# MCA example: Pre- and extramarital sex

- PreSex data: the  $2 \times 2 \times 2 \times 2$  table of gender, premarital sex, extramatrial sex and marital status (divorced, still married)
- The function mjca () provides several scalings for the singular values
- Here I use lambda="Burt"

### MCA example: Pre- and extramarital sex

plot (presex.mca)



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Multiple correspondence analysis MCA inertia Multiple correspondence analysis MCA inertia

#### Inertia in MCA

- In simple CA, total inertia =  $\sum \lambda_i^2 = \chi^2/n$
- sensible to consider % inertia for each dimension

Not so straight-forward in MCA:

- For a given indicator matrix,  $Z_q$ , the inertia is  $J_q 1$
- For all variables, with  $J = \Sigma J_q$  categories, the total inertia of  $Z = [Z_1, \dots, Z_Q]$  is the average of the inertias of the sub-tables

$$inertia(\mathbf{Z}) = \frac{1}{Q} \sum_{q} inertia(\mathbf{Z}_q) = \frac{1}{Q} \sum_{q} (J_q - 1) = \frac{J - Q}{Q}$$

- The average inertia per dimension is therefore 1/Q
- $\implies$  Interpret dimensions with inertia > 1/Q (as in PCA:  $\lambda > 1$ )
- In analysis of the Burt matrix, average inertia is inflated by the diagonal blocks

### Inertia in MCA

Two solutions:

#### Adjusted inertia

- Ignores the diagonal blocks in the Burt matrix
- Calculates adjusted inertia as

$$(\lambda_i^{\star})^2 = \left[\frac{Q}{Q-1}(\lambda_i^Z - \frac{1}{Q})\right]^2$$

• Express contributions of dimensions as  $(\lambda_i^{\star})^2/\sum(\lambda_i^{\star})^2$ , with summation over only dimensions with  $(\lambda^Z)^2 > 1/Q$ .

#### Joint correspondence analysis

- Start with MCA analysis of the Burt matrix
- Replace diagonal blocks with values estimated from that solution
- Repeat until solution converges, improving the fit to off-diagonal blocks

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Multiple correspondence analysis

MCA inertia

Multiple correspondence analysis MCA

MCA inertia

# MCA example: Survival on the Titanic

- Analyze the Titanic data, using mjca()
- The default inertia method is lambda="adjusted"
- Other methods are "indicator", "Burt", "JCA"

```
data(Titanic)
titanic.mca <- mjca(Titanic)
summary(titanic.mca)

##
## Principal inertias (eigenvalues):
##
## dim value % cum% scree plot
## 1 0.067655 76.8 76.8 **************
## 2 0.005386 6.1 82.9 **
## 3 00000000 0.0 82.9
## Total: 0.088118</pre>
```

#### Compare adjusted inertias with other methods:

```
##
## Principal inertias (eigenvalues):
##
## dim value % cum% scree plot
## 1  0.445079 29.7 29.7 ******
## 2  0.305044 20.3 50.0 *****
## 3  0.250006 16.7 66.7 ****
## 4  0.205037 13.7 80.3 ***
## 5  0.178515 11.9 92.2 ***
## 6  0.116318 7.8 100.0 **
## Total: 1.500000 100.0
```

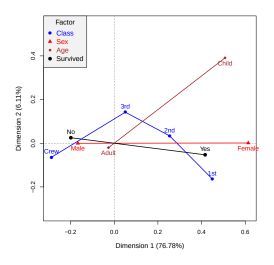
- Total inertia is sum(dim(Titanic)-1)/4 = 6/4 = 1.5
- should only interpret dimensions with inertia > 1/4

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le correspondence analysis MCA inertia Biplots

# MCA example: Survival on the Titanic

plot(titanic.mca)



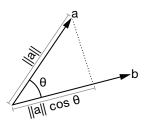
- Dim 1 is perfectly aligned with sex
- This is also strongly aligned with survival and class
- Dim 2 pertains largely to class and age effects
- Survival associated with being female, upper class and child
- Using adjusted inertia, the 2D solution accounts for 83%

### Biplots for contingency tables

The *biplot* is another visualization method that also uses the SVD to give a low-rank (2D) representation.

- In CA, the (weighted)  $\chi^2$  distances between row (column) points reflect the differences among row (column) profiles
- In the biplot, rows and columns are represented by vectors from the origin with an inner product (projection) interpretation

$$\mathbf{Y} \approx \mathbf{A} \mathbf{B}^{\mathsf{T}} \Longleftrightarrow \mathbf{y}_{ii} \approx \mathbf{a}_i^{\mathsf{T}} \mathbf{b}_i$$



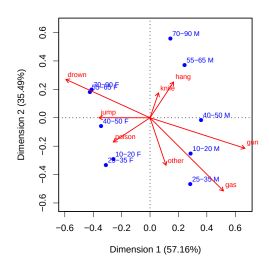
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Summary

Biplots

# Example: suicide rates in Germany

There are a variety of different scalings for biplots. Here we use the contribution biplot



- Associations between age-sex categories and suicide methods can be read as projections of the points on the vectors.
- Lengths of vectors for suicide categories reflect their contributions to this 2D plot

### Summary

- CA is an exploratory method designed to account for association (Pearson  $\chi^2$ ) in a small number of dimensions
  - Row and column scores provide an optimal scaling of the category levels
  - Plots of these can suggest an explanation for association
- CA uses the singular value decomposition to approximate the matrix of residuals from independence
- Standard and principal coordinates have different geometric properties, but are essentially re-scalings of each other
- Multi-way tables can be handled by:
  - Stacking approach— collapse some dimensions interactively to a 2-way table
  - ullet Each way of stacking o a loglinear model
  - MCA analyzes the full n way table using an indicator matrix or the Burt matrix

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