How to build and interpret regression models

Marie Laure Delignette-Muller - VetAgro Sup - LBBE

2022-10-11



Definition of few terms we will use for regression

The dependent variable = the outcome

Independent variables = input variables = regressors = predictors

- ► Independent variable(s) of interest (e.g. the treatment in a clinical trial)
- Covariates = confounding variables or factors = independent variables that may influence the outcome but are not of direct interest (the term factor is used for categorial variables)

Definition of few terms we will use for risk assessment

 $Risk\ factor = risk\ determinant = something\ that\ increases\ the\ chance\ of\ developing\ a\ disease$

Definition found on https://www.cancer.gov/publications/dictionaries/cancer-terms/def/risk-factor.

Implicit **causal interpretation**: alteration of the factor \Rightarrow alteration of the outcome

Potential leverage to reduce the risk if it is controllable.



Risk marker

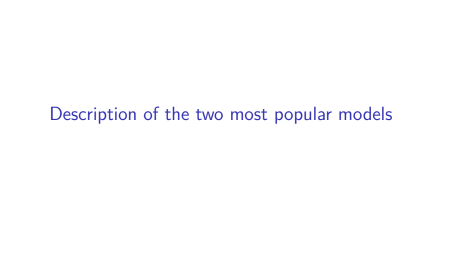
There is an association between the marker and the outcome, but an alteration of the marker will not necessarily affect the outcome.

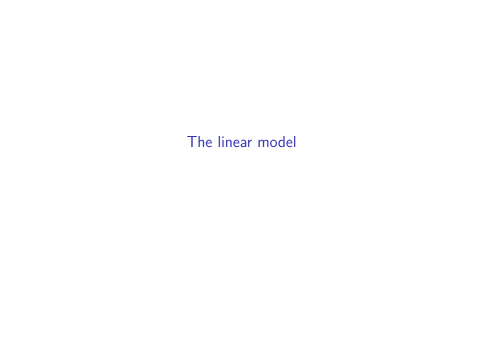
Main preliminar questions to build a model

- ▶ What is the outcome ? The choice of the type of model depends on the nature of the data.
- What are the relevant input variables? We should include the main ones (to avoid confusion bias) but not too many (to avoid a too strong uncertainty on coefficients that would make them useless), and limiting the collinearity between input variables.
- What is the expected relationship between each input and the outcome? Is the linearity assumption reasonable for quantitative input variables?
- ▶ What are the **potential interactions**? Which are the inputs that may have an interaction effect on the outcome?
- What is the purpose of modeling: explicative or predictive? To identify risk factors or risk markers?

Rationale for our presentation plan

- Description of the two most popular of models: their formulation, their hypotheses, the interpretation of their coefficients.
- 2. **Selection of input variables**: why it is necessary, the different strategies proposed and used for the selection of input variables.
- 3. **Model building** in a **risk assessment** perspective: for what purpose ? limits of the approach ?





The linear model: formalization and interpretation

One **continuous outcome** Y and one or more **continuous** and/or **categorial input variables** coded by X_k ,

Each categorial variable with p categories is associated to p-1 dummy variables X_k coding for the membership of each observation to the p groups except the reference one.

$$Y_i = \beta_0 + \beta_1 X_{1i} + \beta_2 X_{2i} + \dots + \beta_k X_{ki} + \dots + \epsilon_i$$

with $\epsilon_i \sim N(0, \sigma)$

Deterministic part: linear link Stochastic part : Gaussian model

Interpretation of the regression coefficients:

- For continuous inputs: β_k estimates the change in the outcome corresponding to a unit change in the input
- For categorial inputs: β_k estimates the difference of the mean in group k to the mean in the reference group

The linear model after log transformation of the outcome

$$In(Y_i) = \beta_0 + \beta_1 X_{1i} + \beta_2 X_{2i} + \dots + \beta_k X_{ki} + \dots + \epsilon_i$$

$$\Leftrightarrow Y_i = e^{\beta_0} \times e^{\beta_1 X_{1i}} \times e^{\beta_2 X_{2i}} \times \dots \times e^{\beta_k X_{ki}} \times \dots \times e^{\epsilon_i}$$

Interpretation of the regression coefficients:

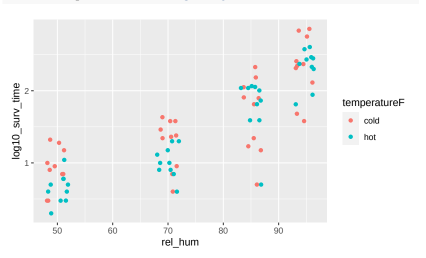
- For continuous inputs: e^{β_k} can be traduced as a multiplicative effect on the outcome corresponding to a unit change in the input
- For categorial inputs: e^{β_k} can be traduced as a multiplicative effect in group k by comparison to the reference group

Go back to our tick example

```
dtot <- read.table("DATA/Milne1950.txt", header = TRUE)</pre>
str(dtot)
## 'data.frame': 100 obs. of 3 variables:
## $ rel hum : int 0 50 70 85 95 0 50 70 85 95 ...
## $ surv time : int 7 7 22 15 38 9 9 23 22 48 ...
   $ temperature: int 5 5 5 5 5 5 5 5 5 5 ...
##
# replacement of 0% humidity by 10%
# as in the paper Wongnak et al. 2022
dtot$rel_hum[dtot$rel_hum == 0] <- 10
# add of the log10 tranformed survival time
dtot$log10_surv_time <- log10(dtot$surv_time)</pre>
dtot$temperatureF <- as.factor(ifelse(dtot$temperature < 15,</pre>
                                       "cold", "hot"))
# Exclusion of the driest condition
dhum <- subset(dtot, rel_hum > 10)
```

Plot of data

```
ggplot(data = dhum, aes(x = rel_hum, y = log10_surv_time,
col = temperatureF)) + geom_jitter(width = 2)
```



Fit of a model with the relative humidity as quantitative and the temperature as a categorial variable

```
(mancova <- lm(log10_surv_time ~ rel_hum + temperatureF,
                 data = dhum))
##
## Call:
## lm(formula = log10_surv_time ~ rel_hum + temperatureF, data = dhum)
##
## Coefficients:
      (Intercept) rel_hum temperatureFhot
##
##
          -0.9533
                          0.0333
                                           -0.1065
# coefficients traduced in multiplicative factors
exp(coef(mancova)[2:3])
##
          rel_hum temperatureFhot
```

0.899

1.034

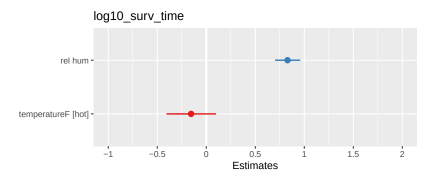
##

Plot of the coefficients as additive effects on Y log scale

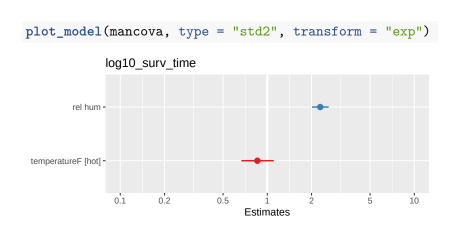
To use the same scale for all the coefficients, the β_k associated to continuous inputs may be multiplied by $2 \times SD(X_k)$ as below.

Interpretation: outcome change for a change of 2 standard deviations of the input.

```
plot_model(mancova, type = "std2")
```



Plot of the coefficients as multiplicative effects on Y raw scale



An example with many continuous and categorial input variables

Pankova *et al.* 2018. **Early weight gain after stopping smoking:** a predictor of overall large weight gain? A single-site retrospective cohort study. BMJ open, 8(12), e023987.

A study based on 1050 patients who stopped smoking with a linear model linking a continuous outcome (relative change in weight 1 year after smoking cessation) with many various input variables.

Try to interpret the reported results

8 Open access Multiple linear model for relative change in weight 1 year after smoking cessation Table 5 Multiple model (n=765) Model with multiple imputation (n=772) Personal characteristics Beta (95% CI) P values Beta (95% CI) P values Weight change in third month (%) 0.134 (-0.037 to 0.304) 0.124 0.141 (-0.027 to 0.31) 0.101 0.804 (0.039 to 1.57) 0.039 0.781 (0.018 to 1.544) 0.045 Female Age at baseline visit (years) -0.005 (-0.034 to 0.024) 0.736 -0.006 (-0.034 to 0.023) 0.682 BMI (ka/m²) -0.209 (-0.3 to -0.118) 0.000 -0.202 (-0.292 to -0.111) < 0.001 BDI score 0.021 (-0.037 to 0.078) 0.467 0.004 (-0.211 to 0.218) 0.974 FTCD score 0 (-0.214 to 0.214) 1.000 Cigarettes per day 0.033 (-0.018 to 0.084) 0.207 0.03 (-0.021 to 0.081) 0.243 Age at regular smoking initiation (years) 0.05 (-0.046 to 0.145) 0.307 0.045 (-0.05 to 0.14) 0.356 Bupropion* 0.011 (-1.337 to 1.359) 0.987 -0.049 (-1.406 to 1.309) 0.944 Varenicline* -0.384 (-1.267 to 0.5) 0.395 -0.349 (-1.225 to 0.527) 0.435 Nicotine replacement therapy* 0.010 -1.068 (-1.879 to -0.257) 0.010 Physical activity Regularly (more times weekly) -1.015 (-2.072 to 0.042) 0.060 -0.957 (-2.01 to 0.095) 0.075 Weekly -0.137 (-1.185 to 0.911) 0.798 -0.1 (-1.146 to 0.946) 0.851 Irregularly 0.402 (-0.657 to 1.462) 0.457 0.466 (-0.586 to 1.519) 0.385 ref. ref. Never Intercept 8.411 (4.993 to 11.828) < 0.001 8.208 (4.795 to 11.621) < 0.001

BDI, Beck Depression Inventory; BMI, body mass index; FTCD, Fagerström Test of Cigarette Dependence.

^{*}Using of specified pharmacotherapy (in monotherapy or in combination with other therapy).

Extracts from the abstract

Results

The regression coefficient per 1% rise in the first 3 months was +0.13% (95% CI -0.04% to 0.30%). In addition, lower body mass index was associated with greater weight gain, while using nicotine replacement therapy was associated with less weight gain at 1-year follow-up.

Conclusions

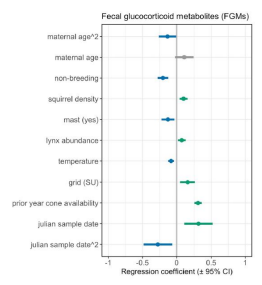
People who stop smoking and gain a larger amount of weight early after quitting are not more likely to gain excessively at 1 year.

What type of plot could help us to interprete those estimations?

Another similar example presented using a forest plot

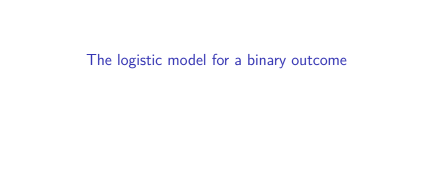
Petrullo *et al.* 2022. The glucocorticoid response to environmental change is not specific to agents of natural selection in wild red squirrels. Hormones and Behavior, 146, 105262.

Another similar example presented using a forest plot - Figure 5



Another similar example presented using a forest plot - Figure 5 caption

Fig. 5. Female red squirrels exhibit a general, rather than specific, glucocorticoid response to environmental change. Forest plot of model estimates and associated confidence intervals corresponding to linear mixed-effects model testing the effects of the same potential ecological agents of selection from Fig. 4 on concentrations of red squirrel fecal glucocorticoid metabolites (FGMs). The dataset contained 1298 FGM measures from 165 females across 7 years, and the model included maternal ID as a random factor (not shown, explained variance in FGM concentrations 0.03). Continuous fixed variables were standardized to a mean of zero and unit variance, and FGMs were log-transformed to achieve residual normality. We controlled for potential effects of intrinsic factors (i.e., maternal (linear and quadratic) age and reproductive status (breeding or non-breeding) on FGM concentrations. Squirrel density, mast (yes/ no), and temperature were assessed the same as in selection models. We included lynx abundance as a continuous variable rather than a categorical variable of lynx-hare cycle as we expected squirrels to exhibit endocrine responses to the presence of lynx regardless of the abundance of hares. Nonsignificant effects are shown in gray; positive effects denoted in green, and negative effects in blue.



The logistic model for a binary outcome

One binary outcome Z and one or more continuous and/or categorial input variables coded by X_k ,

Stochastic part: $Z_i \sim Bernoulli(p_i)$

Deterministic part:

$$logit(p_i) = ln(\frac{p_i}{1-p_i}) = \beta_0 + \beta_1 X_{1i} + \beta_2 X_{2i} + \dots + \beta_k X_{ki} + \dots$$

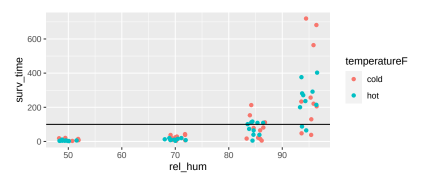
$$\Leftrightarrow \frac{p_i}{1-p_i} = odd_i = e^{\beta_0} \times e^{\beta_1 X_{1i}} \times e^{\beta_2 X_{2i}} \times \dots \times e^{\beta_k X_{ki}} \times \dots$$

Interpretation of the regression coefficients:

- For **continuous** inputs: e^{β_k} can be traduced as a **multiplicative effect on the odd** = odds ratio (OR) corresponding to a **unit change in the input**
- For categorial inputs: e^{β_k} can be traduced as the odds ratio (OR) for group k by comparison to the reference group

Take the example of tick survival by studying the survival or not after 100 days

```
ggplot(data = dhum, aes(x = rel_hum, y = surv_time,
col = temperatureF)) + geom_jitter(width = 2) +
geom_hline(yintercept = 100)
```



Fit of a GLM (logistic regression)

```
dhum$survabove100days <- as.factor(dhum$surv_time > 100)
(mlogis <- glm(survabove100days ~ rel_hum + temperatureF,</pre>
                  family = "binomial", data = dhum))
##
## Call: glm(formula = survabove100days ~ rel_hum + temperatureF, fami
##
      data = dhum)
##
## Coefficients:
       (Intercept) rel_hum temperatureFhot
##
##
          -19.831
                             0.223
                                               0.520
##
## Degrees of Freedom: 79 Total (i.e. Null); 77 Residual
## Null Deviance:
                        97.7
## Residual Deviance: 47.5 AIC: 53.5
```

Estimated coefficients

[1] 1.25 1.68

```
## Coefficients

## Estimate Std. Error z value Pr(>|z|)

## (Intercept) -19.831 5.1383 -3.859 0.000114

## rel_hum 0.223 0.0576 3.866 0.000110

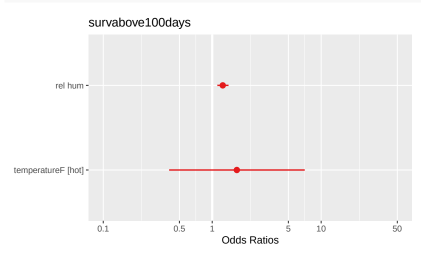
## temperatureFhot 0.520 0.7280 0.715 0.474846

# Effects traduced in odds ratios

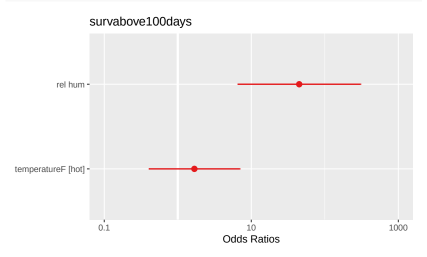
exp(summary(mlogis)$coefficients[2:3])
```

Plot of the coefficients after exponential transformation of them to have ORs

plot_model(mlogis)



Plot of the coefficients after 1/multiplication by $2 \times SD(X_k)$ and 2/exponential transformation



Now look at some logistic regression results in papers

Risk factors associated to leptospirosis

Hinjoy et al. 2019. Environmental and behavioral risk factors for severe leptospirosis in Thailand. Tropical medicine and infectious disease, 4(2), 79. T4

Dung et al. 2022. A case–control study of agricultural and behavioral factors associated with leptospirosis in Vietnam. BMC Infectious Diseases, 22(1), 1-8. T2 and 4

Risk factors associated to HUS on 411 STEC strains collected on patients

De Rauw *et al.* 2019. Risk determinants for the development of typical haemolytic uremic syndrome in Belgium and proposition of a new virulence typing algorithm for Shiga toxin-producing *Escherichia coli*. Epidemiology & Infection, 147. T2

Risk factors associated to STEC positivity of farms

Henry *et al.* 2017. British *Escherichia coli* O157 in Cattle Study (BECS): to determine the prevalence of *E. coli* O157 in herds with cattle destined for the food chain. Epidemiology & Infection, 145(15), 3168-3179. T2

Patterson *et al.* 2022. Risk factors of Shiga toxin-producing *Escherichia coli* in livestock raised on diversified small-scale farms in California. Epidemiology & Infection, 150. T4

Hinjoy et al. 2019 - risk factors associated to leptospirosis

 $\textbf{Table 4.} \ \ \textbf{Unadjusted and adjusted odds ratios and } 95\% \ confidence \ intervals \ risk \ factors \ for \ leptospirosis.$

Risk Factors	Severe Leptospirosis n (%) N = 33	Non-severe Leptospirosis n (%) N = 11	Unadjusted OR (95% CI)	Adjusted OR (95% CI)	
Bathing in natural bodies of	29	5	7.25	10.45	
water 2 weeks before illness	(87.9)	(50.0)	(1.43-36.69)	(1.17 - 93.35)	
Living nearby rubber	18	1	12.00	11.65	
tree plantations	(54.6)	(9.1)	(1.37-104.77)	(1.08-125.53)	
Male	27	7	2-57	3.61	
	(81.8)	(63.6)	(0.57-11.69)	(0.40-32.23)	
Having	11	4	0.88	3.81	
underlying disease(s)	(33.4)	(36.4)	(0.21-3.64)	(0.39 - 37.75)	

Dung et al. 2022 - risk factors associated to leptospirosis

Table 2 Multiple logistic regression analysis of occupations and agricultural practices

Variable	OR (95% CI)	p	
Cultivating	2.83 (1.38–5.79)	0.004	
Animal farming	8.26 (2.24–30.52)	0.002	
Pig owners	10.48 (5.05-21.73)	0.000	
Cat owners	2.62 (1.49-4.61)	0.001	

Dung et al. 2022 - risk factors associated to leptospirosis

Table 4 Multiple logistic regression analysis of behavioral factors

Variable	OR (95% CI)	p
Hand washing after using toilet	0.39 (0.23–0.68)	0.001
Hand washing after farming / gardening	0.57 (0.38-0.86)	0.007
Bathing after farming, gardening, cattle / poultry contact	0.33 (0.19–0.58)	0.000
Drinking unboiled water	1.72 (1.14-2.59)	0.010

De Rauw *et al.* 2019 - risk factors associated to HUS on 411 STEC strains collected on patients

Table 2. Statistically significant results of the multivariate logistic regression models A (*stx1* (with or without *stx2*) *vs. stx2* (with or without *stx1*) positives), B (*stx1* (not *stx2*) positives, *stx1+stx2* positives, *vs. stx2* (not *stx1*) positives) and C (significant *stx1* and *stx2* subtypes)

Statistically significant variable	<i>P</i> value	Odds ratio (95% CI) ^a			
Model A: stx1 (with or without stx2) vs. stx2 (with or without stx1) positives					
Higher risk for HUS development					
Patient age ≤5 years	0.05	7.9 (1.0-62.2)			
Patient age 6–12 years	0.02	13.1 (1.6-107.0)			
Patient age ≽75 years	0.04	10.7 (1.2-95.5)			
stx2 gene (with or without stx1)	0.03	5.4 (1.2-25.1)			
eae gene	0.04	2.8 (1.0-7.7)			
Lower risk for HUS development					
stx1 gene (with or without stx2)	0.00	0.3 (0.1-0.5)			

Henry et al. 2017 - risk factors associated to E. coli O157 positivity of farms

Table 2 Final multivariable logistic model for Outcome 1a for the Scotland data

Variable	Value	Coefficient	OR [95% CI]	<i>p</i> -value
Intercept		-1.04		0.040
breeding females brought on (BFBO)	No	Baseline	1.00	
	Yes	1.71	5.52 [1.81–16.89]	0.003
cattle 12–30 months		0.009	1.009 [1.001-1.02]	0.028
bought other livestock	No	Baseline	1.00	I
	Yes	-1.45	0.23 [0.07-0.75]	0.015
season	Autumn	Baseline	1.00	
	Winter	-0.53	0.59 [0.16-2.20]	0.433
	Spring	-2.33	0.10 [0.02-0.56]	0.009
	Summer	-0.45	0.64 [0.18-2.30]	0.493

OR Odds ratio; CI Confidence Interval

Significant ($p \le 0.05$) OR and p-values are highlighted in bold text McFadden's pseudo $R^2 = 0.21$

^a Farm classified as positive for E. coli O157

Patterson *et al.* 2022 - risk factors associated to STEC positivity of farms

Table 4. Association between risk factors and the presence of Shiga toxin-producing *Escherichia coli* in faecal samples (n = 502) collected from 14³ diversified small-scale farms in California between May 2015 and June 2016, as demonstrated by a generalised linear mixed model, with farm as a random effect

Variable	Variable category	Estimate	OR	95% CI	P-value
Intercept		-2.57			0.001*
Daily maximum temperature °C	numeric	-0.06	0.95	0.91-0.98	0.003*
Sample source species	Swine	Reference			
	Goats	0.97	2.64	0.90-7.70	0.076
	Sheep	1.67	5.29	1.80-15.51	0.002*
	Cattle	1.53	4.61	1.64-12.96	0.004*
Multiple species shared a barn	No	Reference			
	Yes	1.83	6.23	1.84-21.15	0.003*
Livestock were allowed contact with wild areas	No	Reference			
	Yes	1.29	3.63	1.37-9.62	0.009*
Number of years in operation	6–30 years	Reference			
	1–5 years	-0.98	0.38	0.13-1.11	0.076

^aTwo of the 16 participant surveys were not completed.
*Indicates statistical significance with a P-value < 0.05.</p>

Can we say that having multiple species sharing a same barn multiplies by 6.23 the risk for a farm to be STEC positive?

Do not confound odds ratios and risk ratios

Coefficients estimated by **logistic regression** are **odds ratios**

between two conditions
$$k$$
 and 0 (reference): $OR = \frac{\frac{\rho_k}{1-\rho_k}}{\frac{\rho_0}{1-\rho_0}}$

A more intuitive statistics is risk ratio: $RR = \frac{p_k}{p_0}$

Relation between OR and RR: $OR = RR \times \frac{1-p_0}{1-p_k}$

OR is a good approximation of RR only if p_k and p_0 are close to 0. If this is not the case OR exaggerates the RR.

(If
$$p_k > p_0$$
, $RR > 1$ and $1 - p_0 > 1 - p_k$ so $\frac{1 - p_0}{1 - p_k} > 1$ If $p_k < p_0$, $RR < 1$ and $1 - p_0 < 1 - p_k$ so $\frac{1 - p_0}{1 - p_k} < 1$)

BE CAREFUL! Interpretation of ORs as RRs is a very common error.

A reference that shows the impact of this common error

Sheldrick et al. 2017. Math matters: how misinterpretation of odds ratios and risk ratios may influence conclusions. Academic Pediatrics, 17(1), 1-3.

C Degree to which OR overestimates RR [OR/RR]

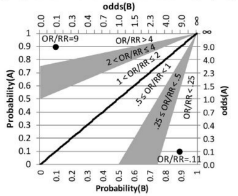
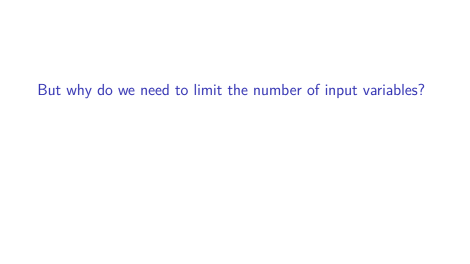


Figure 1. Relationship between RR and OR. (A) RR = probability A/ probability B. (B) OR. (C) Degree to which OR overestimates RR (OR/RR). RR indicates relative risk; OR, odds ratio.

Selection of input variables



Let us look at in-field data collected on Nile monitors

An example from Ciliberti et al. 2011.

Ciliberti et al. 2011. The Nile monitor (Varanus niloticus; Squamata: Varanidae) as a sentinel species for lead and cadmium contamination in sub-Saharan wetlands. Science of the Total Environment, 409(22), 4735-4745.

Nile monitors (large African lizards) were captured in different areas of Africa. The lead content in their kidneys was determined and different morphometric parameters were measured on these animals. We wish to build a **model describing the decimal logarithm of the lead content** (log10Pb) as a function of the variables **sex** (sex), the **area of capture** (site), chosen to represent gradient of contamination level), the **fat somatic index** (FS), the **snout-vent length** (in log_{10} log10SVL) and the **body mass** (in log_{10} log10BM).

Fit the corresponding linear model (neglecting potential interactions) and carefully look at the results.

Importation of the data

##

female 17 5 3 5 ## male 15 9 4 13

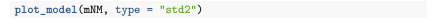
```
dNM <- read.table("DATA/Nilemonitor.txt", header = TRUE,</pre>
                 stringsAsFactors = TRUE)
str(dNM)
## 'data.frame': 71 obs. of 6 variables:
##
   $ sex : Factor w/ 2 levels "female", "male": 2 2 2 2 2 2
   $ site : Factor w/ 4 levels "dif", "fla", "nia", ...: 4 4 4 4
##
   $ log10BM : num -0.108 0.373 0.25 0.334 0.491 ...
##
## $ log10SVL: num 1.51 1.66 1.65 1.67 1.73 ...
## $ FS : num 0.0526 0.0441 0.0826 0.0532 0.0529 ...
   $ log10Pb : num 2.04 1.3 2.14 1.43 2.08 ...
##
xtabs(data = dNM, ~ sex + site)
##
          site
## sex dif fla nia nio
```

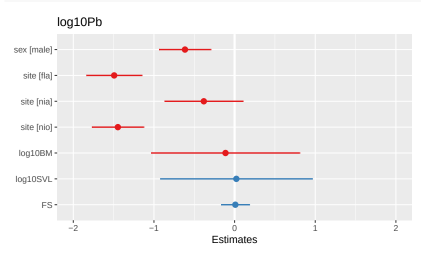
Fit of the model

```
mNM <- lm(log10Pb ~ sex + site + log10BM + log10SVL + FS, data = dNM)
summary(mNM)$coefficients</pre>
```

```
Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 2.5724
                        3.2317 0.7960 4.29e-01
## sexmale -0.2491 0.0651 -3.8256 3.02e-04
## sitefla -0.6045
                       0.0698 -8.6573 2.54e-12
## sitenia -0.1545
                       0.0988 -1.5639 1.23e-01
## sitenio -0.5859 0.0652 -8.9875 6.81e-13
## log10BM -0.1634 0.6681 -0.2445 8.08e-01
## log10SVL 0.0926
                       2.0242 0.0457 9.64e-01
## FS
              0.1603
                        1.5182 0.1056 9.16e-01
```

Forest plot of the coefficients



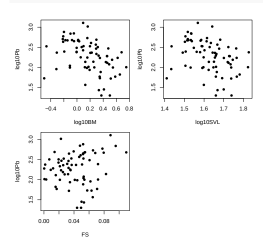


Carefully look at the estimated coefficients for log10BM, log10SVL and FS.

- Do their sign correspond to what is expected ?
- ➤ To answer look at the bivariate correlations between each of those three model inputs and log10Pb.
- ➤ To find an explanation look at the pairwise correlations between those inputs.

Bivariate correlations between log10Pb and the three inputs.

```
par(mfrow = c(2,2)); par(mar = c(4, 4, 1, 1))
plot(log10Pb ~ log10BM, data = dNM, pch = 16)
plot(log10Pb ~ log10SVL, data = dNM, pch = 16)
plot(log10Pb ~ FS, data = dNM, pch = 16)
```



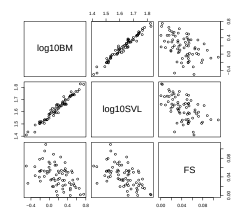
What are the expected signs of the three coefficients?

```
cor(dNM$log10Pb, dNM$log10BM)
## [1] -0.415
cor(dNM$log10Pb, dNM$log10SVL)
## [1] -0.362
cor(dNM$log10Pb, dNM$FS)
## [1] 0.131
  negative for log10BM.
  negative for log10SVL. Why isn't it negative?
  positive for FS.
coef(mNM)[6:8]
```

```
## log10BM log10SVL FS
## -0.1634 0.0926 0.1603
```

Pairwise correlations between those inputs

```
par(mar = c(1, 1, 1, 1)); pairs(dNM[, 3:5])
```



Collinearity between input variables (here log10BM and log10SVL) not only increases the uncertainty of the estimates, but also makes their interpretation meaningless.

But can we use only bivariate analyses?

-0.673 -0.215 -0.677

Comparison of the estimations with and without the sex as an input in the previous example.

```
mNMsexsite <- lm(log10Pb ~ sex + site, data = dNM)
mNMsite <- lm(log10Pb ~ site, data = dNM)
coef(mNMsexsite)[3:5]

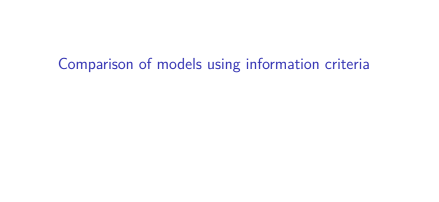
## sitefla sitenia sitenio
## -0.626 -0.187 -0.607
coef(mNMsite)[2:4]

## sitefla sitenia sitenio
```

```
No especially when the design is not balanced (or due to interaction effects, not taken into account here just for simplicity).
```

Conclusion on this rather simple example (in comparison to realistic examples in epidemiology)

- It is important to prevent introducing collinear inputs in a model.
- More generally we need a strategy to choose inputs to include in a model.



Comparison of models using information criteria

- Various strategies are proposed, often based on a predictive perspective.
- Some are based on hypothesis tests for comparing nested models.
- Some are based on information criteria (the most popular being the Akaike's one: AIC). Based on their AIC values, any number of models can be ranked, whatever they are nested or not.

Likelihood and deviance

The **likelihood** is generally expressed in log:

$$logLik = ln(Pr(y \mid \beta, \sigma^2))$$

The **deviance** is the difference in likelihood between the fitted model and the saturated model (perfect model exactly describing the data)

$$Dev = -2logLik(model) + 2logLik(saturated_model)$$

$$Dev = -2logLik(model)$$

if we consider the loglikelihood of the saturated model equal to 1.

Is the model with the smallest deviance always better?

NO

Including new input variables in a model always decreases the deviance,

- but also increases the complexity of the model,
- and so increases the uncertainty on the parameter estimates,
- decreases its robustness to outliers,
- and so decreases its ability to predict new data.

So a compromise must be found: build **parsimonious models**, with just the right number of input variables to well fit the data.

Aikake Information Criterion (AIC)

Information criteria were proposed to help us find a **balance between goodness-of-fit and complexity**, to build parsimonious models.

The most popular one is the **Akaike's information criterion** (AIC) in which the deviance is penalized by twice the number of estimated parameters p:

$$AIC = -2 \times logLik + 2 \times p$$

Given a set of models, the **one with the smaller AIC will be preferred**.

Other popular information criteria

The AIC corrected for small sample size

A correction for sample size (n) is recommended when $\frac{n}{p} < 40$.

$$AICc = -2 \times logLik + 2 \times p + \frac{2p(p+1)}{n-p-1}$$

Bayesian Information Criterion

$$BIC = -2 \times logLik + ln(n) \times p$$

As its penalization for complexity is stronger, it tends to select simpler models than the AIC.

Origin of the AIC

AIC was created to select the best models in a predictive purpose.

The penalization of the deviance in the AIC definition is here just to correct for the overestimation of the error expected in prediction with this model, as the deviance is calculated on the data used for fitting the model (and not on new data - external validation).

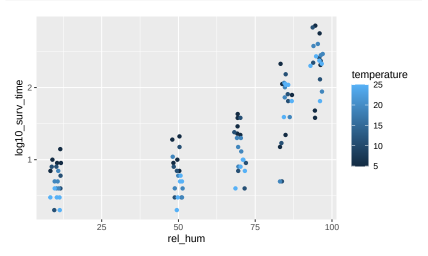
So the challenge of the selection of variables using the AIC is to select the smallest number of input variables that best predicts the outcome.

Comparison of AIC values using R - Ixodes ricinus data

Let us take as an example the *Ixodes ricinus* data (Milne 1950) on which we introduced the linear model.

```
dtot <- read.table("DATA/Milne1950.txt", header = TRUE)</pre>
str(dtot)
## 'data.frame': 100 obs. of 3 variables:
## $ rel hum : int 0 50 70 85 95 0 50 70 85 95 ...
## $ surv time : int 7 7 22 15 38 9 9 23 22 48 ...
   $ temperature: int 5 5 5 5 5 5 5 5 5 5 ...
##
# replacement of 0% humidity by 10%
# as in the paper Wongnak et al. 2022
dtot$rel hum[dtot$rel hum == 0] <- 10
# add of the log10 tranformed survival time
dtot$log10 surv time <- log10(dtot$surv time)</pre>
```

Plot of the data



Comparison of several models

```
# null model
m0 <- lm(log10_surv_time ~ 1, data = dtot)
# linear model
m1 <- lm(log10 surv time ~ rel hum + temperature, data = dtot)
# quadratic model (response surface)
m2 <- lm(log10 surv time ~ rel hum + I(rel hum^2) +
  temperature + I(temperature^2) + rel hum:temperature, data = d
AIC(m0, m1, m2)
## df ATC
## m0 2 214.7
## m1 4 128.7
## m2 7 65.5
```

The more complex model clearly appears as the best one for the prediction of the survival rate. But is it too complex?

Estimates of the most complex model

summary(m2)\$coefficients

```
##
                      Estimate Std. Error t value Pr(>|t|)
  (Intercept)
                      1.501451
                                 2.20e-01
                                           6.831 8.26e-10
## rel hum
                     -0.029433
                                5.08e-03 -5.798 8.93e-08
## I(rel_hum^2)
                                4.35e-05
                      0.000409
                                           9.403 3.42e-15
## temperature
                     -0.049098
                                2.50e-02 -1.963 5.27e-02
## I(temperature^2) 0.000581
                                7.69e-04 0.756 4.52e-01
## rel_hum:temperature 0.000314
                                 1.41e-04
                                           2.228 2.83e-02
```

Is it possible to compare all the possible models?

```
If the number of potential predictors is low (here 5 predictors, rel_hum, temperature, rel_hum^2, temperature^2 and the interaction) it remains possible (here 2^5=32 combinations) but if it is large, it seems difficult (e.g. with 10 potential predictors, 2^{10}=1024 combinations).
```

Stepwise algorithms to choose the best submodel

The R popular function step() proposes three stepwise methods to select the best model based on AIC:

- backward elimination: we start from the most complex considered model, and at each step we remove the predictor that best improves the fit, until the AIC cannot be reduced by removing a predictor.
- forward selection: we start from a minimal model (with the predictors you absolutely want to keep), and at each step we add the predictor that best improves the fit, until the AIC cannot be reduced by adding a predictor.
- both (sometimes called stepwise): a combination of both algorithms, in which at each step predictors can be added or removed, until the AIC cannot be reduced by adding or removing a predictor.

Backward elimination from model m2

```
step(m2, direction = "backward", trace = FALSE)
##
## Call:
## lm(formula = log10_surv_time ~ rel_hum + I(rel_hum^2) + temperature
##
       rel hum:temperature, data = dtot)
##
## Coefficients:
                                                     I(rel_hum^2)
##
           (Intercept)
                                    rel_hum
              1.404443
                                   -0.029433
                                                         0.000409
##
##
           temperature rel_hum:temperature
             -0.031672
                                   0.000314
##
```

One term was eliminated.

Forward selection from model m0 to model m2

```
step(m0, scope = log10_surv_time ~ rel_hum + I(rel_hum^2) +
 temperature + I(temperature^2) + rel_hum:temperature,
       direction = "forward", trace = FALSE)
##
## Call:
## lm(formula = log10_surv_time ~ I(rel_hum^2) + rel_hum + temperature
       rel hum:temperature, data = dtot)
##
##
## Coefficients:
                               I(rel_hum^2)
##
           (Intercept)
                                                         rel_hum
              1.404443
##
                                   0.000409
                                                       -0.029433
##
           temperature rel hum:temperature
             -0.031672
##
                                   0.000314
```

On this example backward elimination and forward selection give the same result. But it is rarely the case when the number of potential predictors is large.

Stepwise selection from model m0 to model m2

```
step(m0, scope = log10_surv_time ~ rel_hum + I(rel_hum^2) +
 temperature + I(temperature^2) + rel_hum:temperature,
       direction = "both", trace = FALSE)
##
## Call:
  lm(formula = log10_surv_time ~ I(rel_hum^2) + rel_hum + temperature
       rel_hum:temperature, data = dtot)
##
##
## Coefficients:
##
           (Intercept)
                               I(rel_hum^2)
                                                         rel hum
##
              1.404443
                                   0.000409
                                                       -0.029433
##
           temperature rel_hum:temperature
             -0.031672
                                   0.000314
##
```

The combination of both methods give the same result on this example.

Your turn to handle the step() function

Take the time to handle the step() function with the three options, carefully looking at the outputs given when the argument trace is fixed to TRUE,

in order to be sure you well understand each algorithm.

Comparison of backward elimination results from model mNM using AIC and BIC criteria

```
mNMAIC <- step(mNM, trace = FALSE)
coef(mNMAIC)[-1]
## sexmale sitefla sitenia sitenio log10BM
## -0.250 -0.605 -0.156 -0.586 -0.138
mNMBIC <- step(mNM, k = log(nrow(dNM)), trace = FALSE)
coef (mNMBTC) [-1]
## sexmale sitefla sitenia sitenia
## -0.273 -0.626 -0.187 -0.607
AIC(mNM, mNMAIC, mNMBIC)
## df AIC
```

mNM 9 -10.8 ## mNMAIC 7 -14.8 ## mNMBIC 6 -14.7

Stepwise methods based on other criteria using R

The three algorithms can be performed based on other criteria than AIC (other information criteria or tests for comparing nested models)

- It is possible to use of step() with the **BIC** by fixing the argument k to ln(n).
- ➤ To use tests comparing nested models (p-values) on Gaussian linear models one can use for example the R package olsrr (with ols_step_forward_p(), ols_step_backward_p() or ols_step_both_p() functions)
- ➤ To use the **AICc** one can use the R package MuMIn: its dredge() function uses AICc by default.

Try to compare backward elimination from model mNM using AIC and BIC criteria

Limits of stepwise selection of input variables

- What criterion to choose ?
- What algorithm to choose (backward elimination, forward selection, both)? Even using one criterion, in the more general case the three approaches do not necessarily propose the same best model.
- ▶ Sometimes the AIC difference between two models is rather small (we often consider that an $\triangle AIC < 2$ is not demonstrative). Is thus such an algorithm relevant ?
- ► The best model may contain non-significant coefficients.
- ► The best model may contain meaningless coefficients.
- The best model regarding the AIC does not necessarily respect the conditions of use that must be carefully checked.
- We must keep in mind that based on the AIC we will select the model that best approximates the data using a minimal number of input variables. But is the model useful in an explanatory perspective?



In "Data analysis using regression and multilevel/hierarchical models" - Gelman & Hill 2006)

Gelman, A., & Hill, J. (2006). Data analysis using regression and multilevel/hierarchical models. Cambridge university press. more than 16000 Google Scholar citations in Oct. 2022

- Include all variables that might be expected to be important in predicting the outcome
- Consider the possibility of gathering some inputs in one predictor, for example calculated as a score from several inputs.
- 3. Consider including **interactions** for inputs having large effects.
- 4. Select the predictors to remove following those rules:
- exclude a predictor if its coefficient is non significant and has not the expected sign.
- think hard about coefficients with significant but unexpected sign.
- generally keep coefficient with expected sign even if non significant.
- always keep a coefficient that is significant and with the expected sign.

In other parts of the book the authors give various methods to check the final model, especially using predictive checking based on data simulation.

In "Applied logistic regression" (Hosmer & Lemeshow 2013)

Hosmer Jr, D. W., Lemeshow, S., & Sturdivant, R. X. (2013). Applied logistic regression (Vol. 398). John Wiley & Sons. more than 71000 Google Scholar citations in Oct. 2022

- 1. Bivariate analyses keep all variables associated with the outcome (p-value > 0.25)
- Multivariate analysis with variables selected in step 1 and all variables of known biological importance. Do not recommend nor stepwise nor best subsets selection of variables: the "analyst must be conscious that such methods can yield a biological implausible model or select irrelevant, or noise, variables".
- 3. For each coefficient, compare its values in Steps 1 and 2, and eliminate predictors for which coefficients are of markedly different orders of magnitude. Then compare the simplified model to the complete model using comparison of nested models. (iterative process). Try to reintroduce in the model each variable not selected in step 1.
- 4. Check the **conditions of use** (linearity for continuous inputs, appropriate categories for discrete variables)
- Check for interactions among the variables in the model, adding each plausible interaction one at a time. And check the goodness-of-fit of the final model

In Bursac et al. 2008 (a highly quoted paper in **medicine**)

Bursac, Z., Gauss, C. H., Williams, D. K., & Hosmer, D. W. (2008). Purposeful selection of variables in logistic regression. Source code for biology and medicine, 3(1), 1-8. 2695 Google Scholar citations in Oct. 2022

- 1. Bivariate analyses keep all variables associated with the outcome (p-value > 0.25)
- 2. **Multivariate analysis** with variables selected in step 1.
- Eliminate predictors that are non significant (p-value > 0.1) and not a confounder (assessed by checking that its elimination does not change the estimation of other coefficients by more than 15% or 20%) (iterative process)
- 4. Try to reintroduce in the model each variable not selected in step 1 and keep it if its contribution is significant (p-value < 0.1 or 0.15). (iterative process only concerning the reintroduction of variables)

In Harrison et al. 2018 (a highly quoted paper in ecology)

Harrison, X. A., Donaldson, L., Correa-Cano, M. E., Evans, J., Fisher, D. N., Goodwin, C. E., . . . & Inger, R. (2018). A brief introduction to mixed effects modelling and multi-model inference in ecology. PeerJ, 6, e4794. 1195 Google Scholar citations in Oct. 2022

- Authors prevent the use of backward selection and hypothesis tests when the number of input variables is large and recommend the ranking of competing models using AIC. For balanced experimental designs with few inputs, they left the open question of which method to use between information criteria and tests.
- Authors recommend "hard thinking about hypotheses" underlying the different competing models instead of selection from all possible subsets, so starting from only a handful of models with biological meaning and without collinear inputs.
- They recommend the ranking of models based on the AIC (with correction if needed AICc) to define a "top model set", taking all the models with a ΔAIC from the best one less than 6. and the elimination of models that are nested models of others in the "top set" as AIC is known to tends toward overly complex models.

The final choice among the remaining models in the top set must be argued by the biologist.

The authors recommend the use of data simulation (powerful but underused tool) to check the final model.

What are the strategies used nowadays by authors to build

models for identifying risk factors?

To have an idea, carefully look at the statitical part of some of the previous examples

De Rauw *et al.* 2019. Risk determinants for the development of typical haemolytic uremic syndrome in Belgium and proposition of a new virulence typing algorithm for Shiga toxin-producing *Escherichia coli*. Epidemiology & Infection, 147.

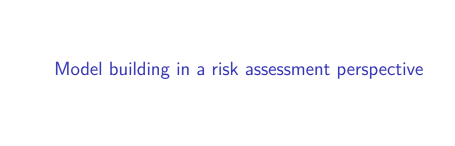
Dung et al. 2022. A case—control study of agricultural and behavioral factors associated with leptospirosis in Vietnam. BMC Infectious Diseases, 22(1), 1-8.

Henry et al. 2017. British Escherichia coli O157 in Cattle Study (BECS): to determine the prevalence of *E. coli* O157 in herds with cattle destined for the food chain. Epidemiology & Infection, 145(15), 3168-3179.

Patterson *et al.* 2022. Risk factors of Shiga toxin-producing *Escherichia coli* in livestock raised on diversified small-scale farms in California. Epidemiology & Infection, 150.

Conclusion of this exploration?

There seems to be
a great variability in used approaches,
and sometimes a gap between
what is recommended and
what is really done.



In which perspective is the model built, inference,

prediction?

In which perspective is the model built, inference, prediction?

- ▶ in a predictive perspective (the one for which information criteria were developed)? We want the model to predict the outcome (e.g. calculate clinical score to do a pronostic)
- in an explicative perspective (inference)? For a better understanding of biological processes? We want to compare models based on different competitive biological hypotheses.

What are the perspectives for **risk management**? What are the main risk factors? (**explicative**) How can I reduce the risk? (**prediction**, using **inputs that are the easiest to control**)

Some references

Tredennick *et al.* 2021. A practical guide to selecting models for exploration, inference, and prediction in ecology. Ecology, 102(6), e03336.

Mac Nally et al. 2018. Model selection using information criteria, but is the "best" model any good? Journal of Applied Ecology, 55(3), 1441-1444.

Go back to Dung et al. 2022

Dung *et al.* 2022. A case–control study of agricultural and behavioral factors associated with leptospirosis in Vietnam. BMC Infectious Diseases, 22(1), 1-8. Tables 3 and 4

Dung et al. 2022 - bivariate analyses

Table 3 Bivariate logistic regression analysis of behavioral factors

Variable	Cases (n = 252)	Controls (n = 252)	OR (95% CI)	p
Hand washing after using tollet	198 (78.6%)	229 (90.9%)	0.37 (0.22-0.62)	0.000
Hand washing after farming/gardening	85 (33.7%)	144 (57.1%)	0.38 (0.27-0.55)	0.000
Hand washing before eating	165 (65.5%)	158 (62.7%)	1.13 (0.78-1.62)	0.516
Hand washing after bathing the livestock or assisting them to breed	93 (36.9%)	1115 (45.6%)	0.69 (0.49-1.00)	0.050
Hand washing after contacting domestic animals	102 (40.5%)	122 (48.4%)	0.73 (0.51-1.03)	0.073
Bathing after farming, gardening, cattle/poultry contact	185 (73.4%)	230 (91.3%)	0.26 (0.16-0.44)	0.000
Using gloves/boots for farming, gardening, livestock/poultry contact	158 (62.7%)	190 (75.4%)	0.55 (0.37-0.81)	0.002
Walking barefoot	171 (67.9%)	171 (67.9%)	-	1
Participating in physical activities	198 (78.6%)	183 (72.6%)	1.38 (0.92-2.01)	0.120
Participating in water sports	21 (8.3%)	30 (11.9%)	0.67 (0.37-1.21)	0.184
Drinking unboiled water	102 (40.5%)	62 (24.6%)	2.08 (1.42-3.05)	0.000
Eating uncooked food	57 (22.6%)	54 (21.4%)	1.07 (0.70-1.63)	0.747

Dung et al. 2022 - multivarite analysis

After backward elimination based on p_values < 0.05.

Table 4 Multiple logistic regression analysis of behavioral factors

Variable	OR (95% CI)	р
Hand washing after using toilet	0.39 (0.23–0.68)	0.001
Hand washing after farming / gardening	0.57 (0.38-0.86)	0.007
Bathing after farming, gardening, cattle / poultry contact	0.33 (0.19–0.58)	0.000
Drinking unboiled water	1.72 (1.14–2.59)	0.010

Dung et al. 2022 - extracted from the paper :

"The multiple logistic regression analysis of behavior risk factors indicated three protective factors,

- hand washing after using toilet,
- hand washing after farming/gardering,
- contacting with cattle and poultry."

What comments/ideas does this example inspire in you? What alternatives could you propose to make the analysis of those data more convincing?

Is it easy to draw conclusions from models fitted on

non-experimental data?

Is it easy to draw conclusions from models fitted on non-experimental data ?

Christenfeld *et al.* 2004. **Risk factors, confounding, and the illusion of statistical control**. Psychosomatic medicine, 66(6), 868-875.

Westfall & Yarkoni, 2016. **Statistically controlling for confounding constructs is harder than you think**. PloS one, 11(3), e0152719.

A realistic "toy" example

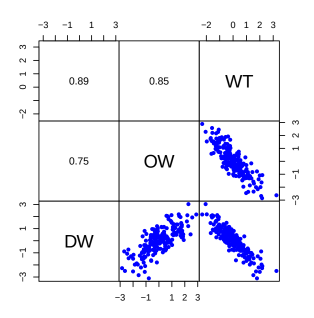
- ▶ **Dog weight** (standardized = centered and reduced): DW
- Owner's weight (standardized): OW
- Common daily walk time (standardized): WT

With $WT \longrightarrow DW$ and $WT \longrightarrow OW$ as causal relationships

Simulation of data from

- \blacktriangleright WT \sim N(0,1)
- \triangleright $OW \sim N(\alpha_{OW} + \beta_{OW} \times WT, \sigma_{OW})$
- $ightharpoonup DW \sim N(\alpha_{DW} + \beta_{DW} \times WT, \sigma_{DW})$

Visualization of simulated data



Linear model: DW as a function of OW

```
mOW <- lm(DW ~ OW)
summary(mOW)$coefficients</pre>
```

```
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.000514 0.0507 0.0101 9.92e-01
## OW 0.707517 0.0449 15.7572 8.64e-37
```

Of course, if we do not take into account WT in the model, we highlight a correlation between DW and OW related to the common causal factor WT.

Linear model: DW as a function of OW and the confounding factor WT

```
mOWWT <- lm(DW ~ OW + WT)
summary(mOWWT)$coefficients
## Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept) 0.0168 0.0343 0.491 6.24e-01

## OW -0.0422 0.0574 -0.735 4.63e-01

## WT -1.0584 0.0688 -15.376 1.43e-35
```

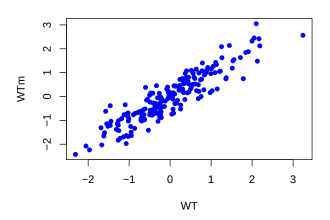
Taking into account the confounding factor WT, as expected, we no longer show a significant effect of OW on DW.

So it works?

Yes, but would it work in real life?

But what if the available WT measurement is noisy (realistic in real life)?

 $WTm \sim N(WT, \sigma_{WTm})$



Linear model: DW as a function of OW and WTm

```
mOWWTm <- lm(DW ~ OW + WTm)
summary(mOWWTm)$coefficients</pre>
```

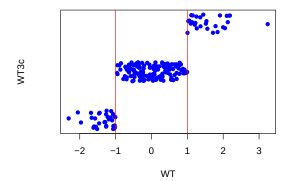
```
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.00398 0.0419 0.0949 9.24e-01
## OW 0.29128 0.0571 5.1014 7.90e-07
## WTm -0.61306 0.0639 -9.6003 3.80e-18
```

Even taking into account the confounding factor WTm (measured with some error), we show a significant effect of OW on DW.

Another realistic case, if only a qualitative measure of WT is used (e.g. categorized in 3 classes)

WT transformed into a categorial variable (WT3c) with three classes:

]-4; -1],]-1; 1],]1; 4].



Linear model: DW as a function of OW and WT3c

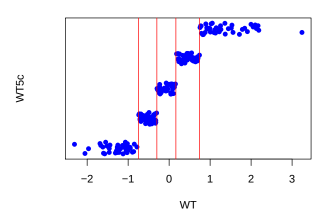
```
mOWWT3c <- lm(DW ~ OW + WT3c)
summary(mOWWT3c)$coefficients</pre>
```

```
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.942 0.1366 6.90 7.08e-11
## OW 0.324 0.0567 5.72 3.86e-08
## WT3c(-1,1] -0.909 0.1463 -6.21 3.05e-09
## WT3c(1,4] -2.119 0.2317 -9.14 7.76e-17
```

We still show a significant effect of OW on DW!

And if the discretization of WT is less coarse and balanced

WT transformed into a categorail variable (WT5c) with five balanced classes, whose limits are defined by the quintiles.



Linear model: DW as a function of OW and WT5c

```
mOWWT5c <- lm(DW ~ OW + WT5c)
summary(mOWWT5c)$coefficients</pre>
```

```
##
                    Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                       1.109
                                0.1247
                                         8.90 4.08e-16
                                0.0613
## NW
                       0.141
                                         2.31 2.21e-02
                                0.1391
## WT5c(-0.75,-0.299]
                      -0.678
                                        -4.87 2.31e-06
## WT5c(-0.299,0.165]
                      -0.985
                                0.1506
                                        -6.54 5.22e-10
## WT5c(0.165,0.743]
                      -1.446
                                0.1687
                                        -8.57 3.26e-15
## WT5c(0.743.4]
                      -2.340
                                0.2121
                                        -11.03 2.83e-22
```

We still show a significant effect of OW on DW!

Conclusion about the possibility of taking into account confounding factors in a linear model

A problem of this kind is very realistic and I let you imagine the consequences!

Taking into account the potential confounding variables in a linear model is essential, but great caution is required when interpreting the results of a linear model on observational data (i.e. with uncontrolled input variables).

Conclusion

Conclusion

- Statistical modeling is a powerful but not perfect tool and should be handled with great caution.
- There is no unique / best strategy to build a model. Authors should be able to well describe and argue their own strategy in order to convince the reader it is well-founded.
- The question of the end use of a model (explicative / predictive) is an underlying question that we should keep in mind while developing any type of model (also crucial in machine learning).