

# t1m User's Guide: Effects under linear, logistic and Poisson regression models with transformed variables

Jose Barrera-Gómez<sup>\*1,2,3</sup> and Xavier Basagaña<sup>†1,2,3</sup>

<sup>1</sup>*Centre for Research in Environmental Epidemiology (CREAL)*

<sup>2</sup>*Universitat Pompeu Fabra (UPF)*

<sup>3</sup>*CIBER Epidemiología y Salud Pública (CIBERESP)*

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## 1 Introduction

This document is a user's guide for the R<sup>1</sup> package `t1m`, which provides the effect of an explanatory variable of interest,  $X$ , on a response variable,  $Y$ , under a linear, logistic or Poisson regression model with transformations in  $X$  and/or  $Y$ . In the case of the linear regression model, log and power transformations in any of  $X$  and  $Y$  are allowed. In the case of logistic and Poisson regression models, log or power transformation in  $X$  are allowed. Other explanatory variables can be in the model, in which case adjusted measures for  $Y$  and adjusted effects of  $X$  on  $Y$  are automatically computed. The package also works if there

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\*jbarrera@creal.cat

†xbasagana@creal.cat

<sup>1</sup>R is a free and open source software and it is available at <http://cran.r-project.org/>.

is no transformation. The package provides both numerical and graphical outputs as well as information on interpreting results. The methodology is described in the manuscript [1], whose illustrative examples, among others, are reproduced here.

## 2 Getting started

Start an R session and load the package by typing

```
> library(tlm)
```

If needed, this user's guide can be recovered by typing

```
> vignette("tlm")
```

You can get a brief overview of the package by typing

```
> help(package = "tlm")
```

### 2.1 Fitting the model

The first step is to fit the model in the transformed space (i.e., considering that  $Y$  and  $X$  are already transformed, if any transformation is assumed). This is performed by the function `tlm`. We can get information on this function using the help:

```
> ?tlm
```

The main arguments of the function `tlm` are:

- **y**: the response variable.
- **x**: the explanatory variable of interest.
- **z**: additional explanatory variables in the model. The default value is **"none"**. For a single variable, we need to write just the name of the variable (unquoted); for more than one variable, the names should be passed unquoted and separated by the plus sign (e.g., `z = x2 + x3 + x4`); in the case of including the remaining variables in **data**, use just a period (`.`). No interaction terms are allowed.
- **family**: the model family. It can be **gaussian**, for linear regression (default); **binomial**, for logistic regression with logit link, or **poisson**, for Poisson regression.
- **data**: a **data.frame** containing the variables **y**, **x** and those in **z**, if any.
- **ypow** and **xpow**: the power transformations assumed in **y** and **x**, respectively. The value 1 (default) means no transformation; the value 0 means log transformation.

We can use the method `summary` to obtain information on the fitted model. As a result of a call to the function `tlm`, we obtain an object of class **"tlm"** which can be passed to the functions `MY` and `effect` in order to obtain adjusted measures and effects, respectively, as we will see in next sections.

## 2.2 Adjusted measures

Once the model has been fitted with the function `tlm`, the resulting object can be passed to the function `MY`, which provides measures of the response variable for given values of the explanatory variable. If the model contains other explanatory variables, then adjusted measures are automatically computed. These adjusted measures are obtained by setting the remaining explanatory variables in the model at their means. We can get information on this function using the help:

```
> ?MY
```

The main arguments of the function `MY` are:

- **object**: an object of class `"tlm"`, the result of a call to the function `tlm`.
- **x**: the value(s) of the explanatory variable of interest for which the expected measure of the response variable should be computed.
- **npoints**: if **x** is not provided, the number of points where the measure should be computed. Default is 10.
- **space**: the space of the variables in which measures should be computed. It can be `"original"` (default) or `"transformed"`.
- **level**: the confidence level for measures. Default is 0.95.

The function `MY` automatically provides the unit of the measure (mean, geometric mean, median, probability or logodds, depending on the case).

In addition, the method `plot` provides, for an object of class `"tlm"`, a graphical representation of adjusted measures. The argument `type` of the method `plot` controls whether the plot should be shown in the original space (`"original"`, default) or in the transformed space (`"transformed"`). The method `plot` automatically labels the  $Y$  axis with the appropriate names of the measure. Using the value `"space = diagnosis"` provides a diagnostics plot.

## 2.3 Adjusted effects

The fitted model can be passed to the function `effect` in order to obtain the effect of  $X$  on  $Y$  in the original space of the variables, adjusted for those variables in  $\mathbf{z}$ , if any. We can get information on this function using the help:

```
> ?effect
```

The main arguments of the function `effect` are:

- **object**: an object of class `"tlm"`, the result of a call to the function `tlm`.
- **x1**: the value(s) of the explanatory variable where the effect should be computed.
- **x2**: the alternative value(s) of the explanatory variable (changing from **x1**) for which the effect should be computed.
- **c**: the additive change in the explanatory variable.

- **q**: the multiplicative change in the explanatory variable.
- **r**: the percent change in the explanatory variable.
- **npoints**: the number of points where the effect should be computed.
- **level**: the confidence level for the effect.

In order to compute effects, the change in the explanatory variable should be characterized. It can be done in several ways. For instance, providing: (1) **x1** and **x2**; (2) **x1** and one of **c**, **q** or **r**; (3) **x1**, **npoints** and one of **c**, **q** or **r**. Only one of the arguments **c**, **q** or **r** is used, prevailing **c** and then **q**. If no enough arguments are passed, the interquartile range will be considered and a summary effect should be computed, if it exists. If the explanatory variable is categorical, then the effect is computing for a change between the reference level and each of the remaining levels of the explanatory variable. Confidence intervals are computed by transforming the endpoints of the intervals in the transformed scale when it is possible, while non-parametric bootstrap is used otherwise. The function **effect** automatically provides the unit of the effect measure (mean, geometric mean, median or odds ratio, depending on the case).

In addition, the function **effectInfo** provides information on interpreting the effect (use **?effectInfo** for further details).

## 3 Illustrative examples

### 3.1 Linear regression model

#### 3.1.1 Log transformation in the response

Consider the evaluation of the association between the intima media thickness of the carotid artery (IMT), measured in mm,  $Y$ , and age, in years,  $X$ . Variable  $Y$  was log transformed to achieve normality. The **imt** data were simulated to emulate true data pattern observed in a real study [2].

First, we can load data and see its first rows and a summary:

```
> data(imt)
> dim(imt)

[1] 2784    3

> head(imt)

  age      imt      logimt
1  53 0.6588690 -0.41723048
2  67 0.9215614 -0.08168583
3  57 0.9539836 -0.04710877
4  55 0.7226028 -0.32489555
5  73 0.7406030 -0.30029063
6  57 0.7638349 -0.26940367

> summary(imt)
```

age	imt	logimt
Min. :32.00	Min. :0.3755	Min. :-0.9794
1st Qu.:51.00	1st Qu.:0.6307	1st Qu.: -0.4609
Median :59.00	Median :0.7257	Median :-0.3206
Mean :58.48	Mean :0.7420	Mean :-0.3196
3rd Qu.:66.00	3rd Qu.:0.8324	3rd Qu.: -0.1834
Max. :86.00	Max. :1.5788	Max. : 0.4567

Further information about data is available with the help function, `?imt`.

Suppose that we are interested in the effect of age on IMT, under a linear regression model with log transformation in the response variable, IMT. The model can be fitted as follows:

```
> modimt <- tlm(y = logimt, x = age, data = imt, ypow = 0)
```

where `ypow = 0` indicates that the response variable is already log transformed. Note that we have not set the family since default family is Gaussian. The fitted model results in:

```
> modimt
```

Linear regression fitted model in the transformed space

-----

Transformations:

In the response variable: log

Call:

```
lm(formula = logimt ~ age, data = imt)
```

Coefficients:

(Intercept)	age
-0.877692	0.009543

Further information on the fitted model is available using the method `summary`:

```
> summary(modimt)
```

Linear regression fitted model in the transformed space

-----

Transformations:

In the response variable: log

Call:

```
lm(formula = logimt ~ age, data = imt)
```

Residuals:

Min	1Q	Median	3Q	Max
-0.57891	-0.11792	0.00142	0.11879	0.61968

Coefficients:

```
      Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.8776919  0.0183979  -47.71  <2e-16 ***
age          0.0095433  0.0003093   30.86  <2e-16 ***
```

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 0.1778 on 2782 degrees of freedom  
Multiple R-squared: 0.255, Adjusted R-squared: 0.2547  
F-statistic: 952.2 on 1 and 2782 DF, p-value: < 2.2e-16

A numerical representation of the relationship between age and IMT is available using the function MY. As default, the measure of the response variable is computed in 10 points along the range of the explanatory variable:

```
> MY(modimt)
```

Estimated adjusted geometric mean of the response variable in the original space:

	X	geomMean(Y)	lower95%	upper95%
1	32	0.5642233	0.5545100	0.5741068
2	38	0.5974736	0.5891267	0.6059388
3	44	0.6326834	0.6257680	0.6396752
4	50	0.6699681	0.6643811	0.6756020
5	56	0.7094501	0.7046577	0.7142750
6	62	0.7512587	0.7460592	0.7564945
7	68	0.7955312	0.7885807	0.8025430
8	74	0.8424128	0.8327802	0.8521567
9	80	0.8920571	0.8791028	0.9052023
10	86	0.9446270	0.9278224	0.9617360

The number of points can be set using npoints:

```
> MY(modimt, npoints = 3)
```

Estimated adjusted geometric mean of the response variable in the original space:

	X	geomMean(Y)	lower95%	upper95%
1	32	0.5642233	0.5545100	0.5741068
2	59	0.7300552	0.7252407	0.7349016
3	86	0.9446270	0.9278224	0.9617360

We can also choose a specific set of values of the explanatory variable for which the measure of the response variable should be computed. For instance, the first and third quartile:

```
> q13 <- quantile(imt$age, probs = c(1, 3)/4)
> MY(modimt, x = q13)
```

Estimated adjusted geometric mean of the response variable in the original space:

```
X geomMean(Y) lower95% upper95%
1 51 0.6763924 0.6709922 0.6818362
2 66 0.7804912 0.7742493 0.7867834
```

Measures can also be computed in the transformed space:

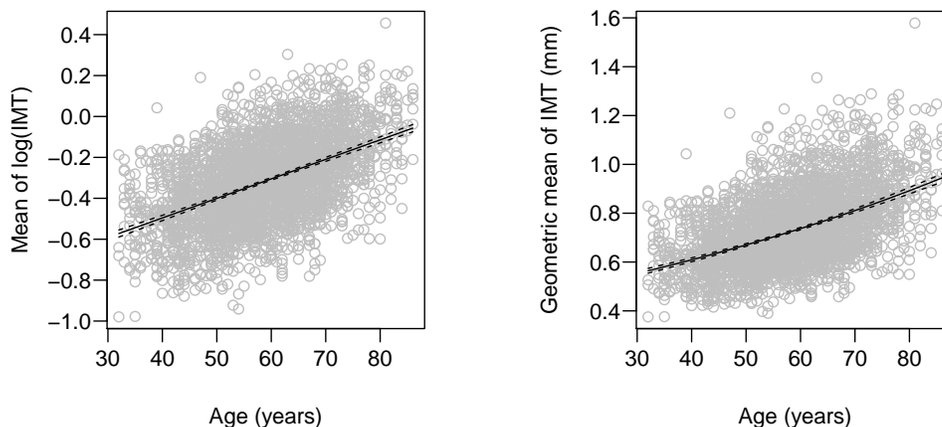
```
> MY(modimt, x = q13, space = "transformed")
```

Estimated adjusted mean of the response variable in the transformed space:

```
X mean(log(Y)) lower95% upper95%
1 51 -0.3909818 -0.3989978 -0.3829659
2 66 -0.2478318 -0.2558614 -0.2398023
```

A graphical representation of the relationship between age and IMT is available using the method `plot`. For instance, the two following instructions provide left and right plots in Figure 1, respectively:

```
> plot(modimt, type = "transformed", observed = T, xname = "Age (years)", yname = "IMT")
> plot(modimt, observed = T, xname = "Age (years)", yname = "IMT (mm)")
```



**Figure 1.** Mean of log of IMT (left) and geometric mean (or median) of IMT (right) as a function of the age. Dashed lines represent 95% confidence intervals for the measure.

The argument `observed` controls whether observations are shown in the plot (default is `FALSE`). Further information on the usage of the method `plot` is available using `?t1m`.

Diagnostics plot as in Figure 2 can be obtained with the following instruction:

```
> plot(modimt, type = "diagnosis")
```

The function `effectInfo` provides information on interpreting the relationship between age ( $X$ ) and IMT ( $Y$ ):

```
> effectInfo(modimt)
```

The effect of  $X$  on  $Y$  can be summarized with a single number as follows:

- Change in  $X$ : additive of  $c$  units
- Type of effect on  $Y$ : percent change in the geometric mean of  $Y$
- Effect size:  $100 * [\exp(c * \beta) - 1]\%$

beta coefficient estimate:

	Estimate	Std. Error	t value	Pr(> t )
age	0.009543334	0.0003092751	30.8571	4.531758e-180

Further details can be obtained using `effect()`.

Thus, we can see that, in this case, if we use additive changes in  $X$  and percent (or multiplicative) changes in the geometric mean of  $Y$ , a summary effect can be obtained, which is independent of the value of  $X$  for which the effect is computed.

The function `effect` provides as default the expected change in IMT for an additive change in age equal to the interquartile range:

```
> effect(modimt)
```

Computing effects...

Adjusted percent change in the geometric mean of the response variable for a 'c' units additive change in the explanatory variable equivalent to the interquartile range:

	c	Estimate	lower95%	upper95%
1	15	15.39029	14.3454	16.44472

For further information on interpreting the effect use `effectInfo()`.

Other measures of effects can be obtained. For instance, we may be interested in a difference of (geometric) means when changing age across the first quartile, the median and the third quartile:

```
> q123 <- quantile(imt$age, probs = 1:3/4) # quartiles  
> effect(modimt, x1 = q123)
```

Computing effects...

Adjusted change in the geometric mean of the response variable when the explanatory variable changes from  $x_1$  to  $x_2$  (confidence interval for the

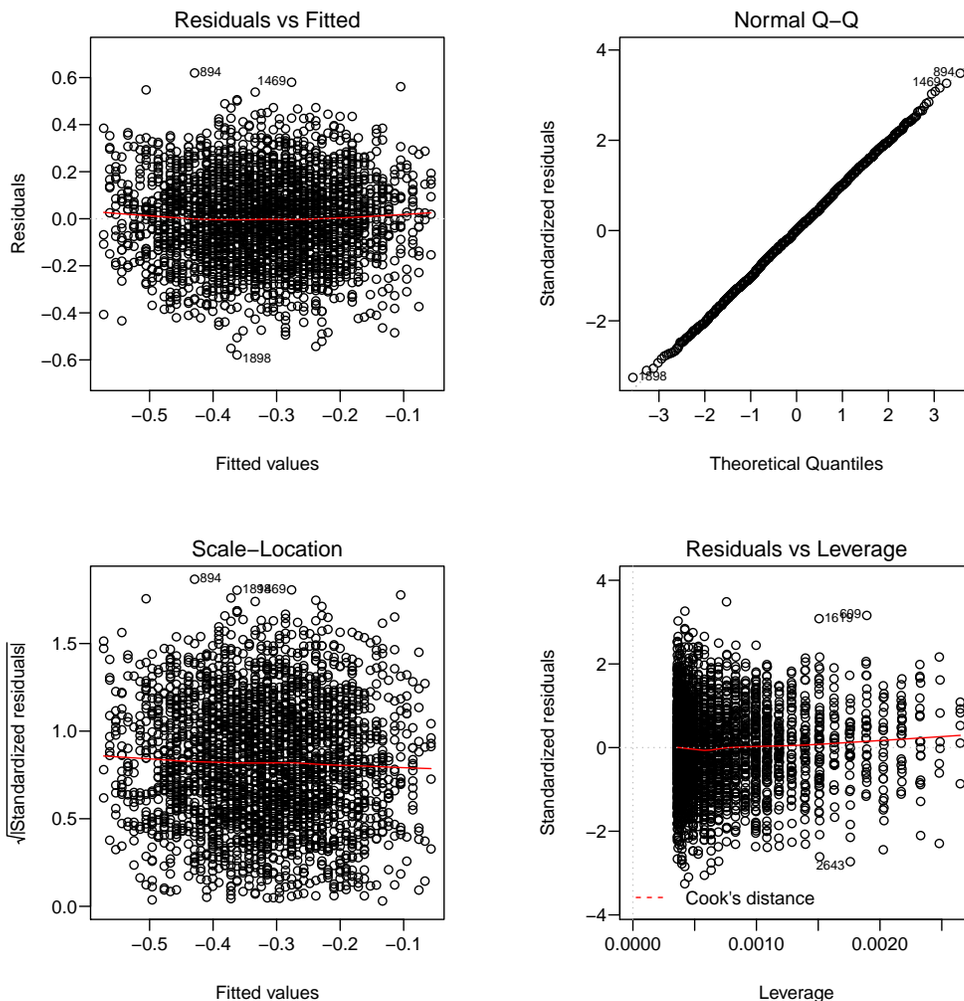


Figure 2. Diagnostics plot for the model modimt.

difference change based on 999 bootstrap samples):

	x1	x2	EstimateDiff	lower95%	upper95%	EstimatePercent	lower95%	upper95%
25%	51	59	0.05366274	0.05039149	0.05674395	7.933669	7.411302	8.458576
50%	59	66	0.05043602	0.04725599	0.05377603	6.908521	6.455654	7.363315

For further information on interpreting the effect use `effectInfo()`.

As in this example, when a summary effect is not computed, then both difference and percent changes in the response are computed. The number of bootstrap samples is controlled by the argument `nboot` whose default value is 999 [3].

### 3.1.2 Log transformation in the explanatory variable

Consider now the evaluation of the association between the birth weight, in grams,  $Y$ , and the cord serum cotinine level in the mother, in ng/ml,  $X$ . Variable  $X$  was log transformed to achieve a more homogeneous distribution. The `cotinine` data were simulated to emulate true data patterns observed in a real study [4]. Data also contains a binary variable indicating whether the birth weight was low (defined as lower than 2500 g).

We can load and explore data (further information about data is available with the help function, `?cotinine`):

```
> data(cotinine)
> dim(cotinine)

[1] 351  4

> head(cotinine)

   cotinine logcotinine weight underweight
2  5.1584035   1.640627  3626          no
5  0.2909473  -1.234613  3672          no
11 4.1119142   1.413889  3779          no
12 3.0037959   1.099877  3540          no
14 5.9240779   1.779025  3179          no
17 7.3854370   1.999510  2494          yes

> summary(cotinine)

   cotinine      logcotinine      weight      underweight
Min.   : 0.200   Min.   :-1.609   Min.   :1656   no :327
1st Qu.: 3.234   1st Qu.: 1.174   1st Qu.:2920   yes: 24
Median : 7.385   Median : 2.000   Median :3232
Mean   :39.431   Mean    : 2.353   Mean    :3218
3rd Qu.:39.226   3rd Qu.: 3.669   3rd Qu.:3522
Max.   :910.000   Max.    : 6.813   Max.    :4771
```

Suppose that we are interested in the effect of cotinine level on birth weight, under a linear regression model with log transformation in the explanatory variable, cotinine. The model can be fitted as follows:

```
> modcot <- tlm(y = weight, x = logcotinine, data = cotinine, xpow = 0)
```

where `xpow = 0` indicates that the explanatory variable is already log transformed. The fitted model provides the following results:

```
> summary(modcot)
```

Linear regression fitted model in the transformed space

-----  
Transformations:

In the explanatory variable: log

Call:

```
lm(formula = weight ~ logcotinine, data = cotinine)
```

Residuals:

Min	1Q	Median	3Q	Max
-1390.64	-280.11	-2.95	300.47	1422.31

Coefficients:

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	3406.49	42.59	79.978	< 2e-16 ***
logcotinine	-80.00	14.95	-5.351	1.58e-07 ***

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 450.1 on 349 degrees of freedom

Multiple R-squared: 0.07583, Adjusted R-squared: 0.07318

F-statistic: 28.64 on 1 and 349 DF, p-value: 1.585e-07

In Figure 3 (obtained with the following instructions) we can see the fitted model (left) and the relationship between cotinine level and birth weight, under the model, in the original space of the variables (right).

```
> plot(modcot, type = "transformed", observed = T, xname = "Cotinine", yname = "weight (kg)")  
> plot(modcot, xname = "Cotinine (ng/ml)", yname = "weight (kg)")
```

The function `effectInfo` provides information on interpreting the relationship between cotinine levels ( $X$ ) and weight ( $Y$ ):

```
> effectInfo(modcot)
```

The effect of  $X$  on  $Y$  can be summarized with a single number as follows:

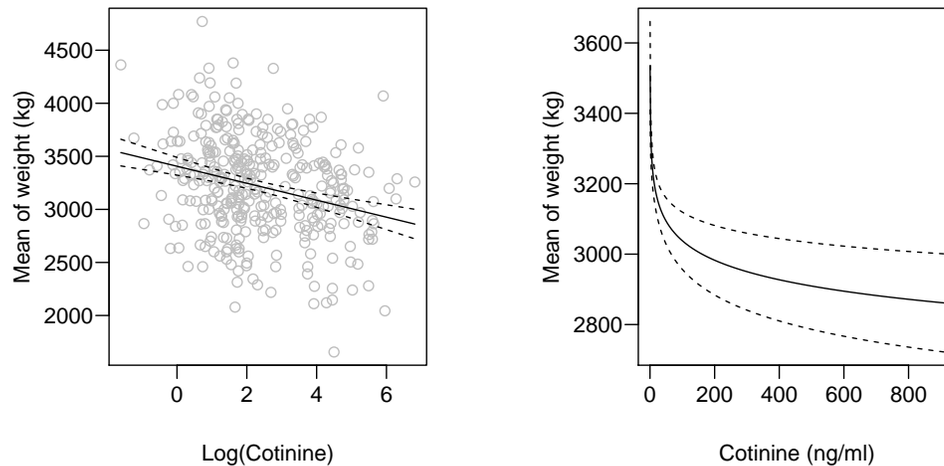
- Change in  $X$ : multiplicative of factor  $q$  (equivalently, adding an  $r = 100 * (q - 1)\%$  to  $X$ )
- Type of effect on  $Y$ : additive change in the mean of  $Y$
- Effect size:  $\beta * \log(q)$  units of  $Y$

beta coefficient estimate:

	Estimate	Std. Error	t value	Pr(> t )
logcotinine	-80.00108	14.94986	-5.351292	1.584903e-07

Further details can be obtained using `effect()`.

In this case, if we use multiplicative (or percent) changes in  $X$  and additive changes in the mean of  $Y$ , a summary effect can be obtained, which is independent of the value of  $X$  for which the effect is computed. The function `effect` provides as default the expected change in weight for a percent change in cotinine levels equal to the interquartile ratio:



**Figure 3.** Mean of birth weight as a function of cotinine in log scale (left) and in the natural scale (right). Dashed lines represent 95% confidence intervals for the measure.

```
> effect(modcot)
```

Computing effects...

Adjusted additive change in the mean of the response variable for an 'r'% change in the explanatory variable equivalent to the interquartile ratio:

```
      r Estimate lower95% upper95%
1 1112.878 -199.6492 -273.0272 -126.2712
```

For further information on interpreting the effect use `effectInfo()`.

Alternatively, by exploring Figure 3, we can see that several 10-fold changes occur in the population and choose the more common number  $q = 10$ , in which case the effect is

```
> effect(modcot, q = 10)
```

Computing effects...

Adjusted additive change in the mean of the response variable for an 'r' = 900% change in the explanatory variable:

```

      r Estimate lower95% upper95%
1 900 -184.2093 -251.9126 -116.506

```

For further information on interpreting the effect use `effectInfo()`.

If we are interested in explore the effect of an additive change in cotinine levels, we can obtain, for example, effects for additive changes in  $X$  along its range. For instance:

```

> range(cotinine$cotinine)

[1] 0.2 910.0

> effect(modcot, x1 = 100, c = 200, npoints = 4)

```

Computing effects...

Adjusted change in the mean of the response variable when the explanatory variable changes from  $x_1$  to  $x_2$  (confidence interval for the percent change based on 999 bootstrap samples):

	x1	x2	EstimateDiff	lower95%	upper95%	EstimatePercent	lower95%	upper95%
1	100	300	-87.89017	-120.19284	-55.58750	-2.8929637	-4.0227274	-1.7692666
2	300	500	-40.86660	-55.88649	-25.84671	-1.3852256	-1.9861551	-0.8069561
3	500	700	-26.91814	-36.81149	-17.02479	-0.9252415	-1.2979132	-0.5732388
4	700	900	-20.10543	-27.49486	-12.71599	-0.6975258	-0.9934569	-0.4388765

For further information on interpreting the effect use `effectInfo()`.

### 3.1.3 Log transformation in both the response and the explanatory variable

Consider an epidemiological study to assess the association between cat allergen levels (*Fel d 1*) in the bed mattress,  $X$ , and in the living room,  $Y$ , in homes of study participants, taking into account cat ownership,  $C$ . Both variables  $X$  and  $Y$  were log transformed to achieve linearity in their relationship. The `feld1` data were simulated to emulate true data patterns observed in a real study [5].

We can load and explore data (further information about data is available with the help function, `?feld1`):

```

> data(feld1)
> dim(feld1)

[1] 471 5

> head(feld1)

      mattress      room logmattress      logroom cat
1 0.66504894 0.26758221 -0.4078946 -1.3183284 yes
2 0.02723504 0.16796992 -3.6032509 -1.7839704 no

```

```

3 0.16773827 0.76728932 -1.7853504 -0.2648913 no
4 0.01391101 0.05480368 -4.2750750 -2.9039979 no
5 0.04216982 0.11931900 -3.1660504 -2.1259547 yes
6 1.44212520 17.29388484 0.3661179 2.8503530 yes

```

```
> summary(feld1)
```

mattress		room		logmattress		logroom		cat	
Min.	: 0.0030	Min.	: 0.0017	Min.	:-5.7975	Min.	:-6.3565	no	:403
1st Qu.:	0.0636	1st Qu.:	0.1076	1st Qu.:	-2.7559	1st Qu.:	-2.2295	yes:	68
Median	: 0.1773	Median	: 0.3081	Median	:-1.7297	Median	:-1.1774		
Mean	: 3.1976	Mean	: 27.5263	Mean	:-1.5942	Mean	:-0.8528		
3rd Qu.:	0.5565	3rd Qu.:	1.1713	3rd Qu.:	-0.5861	3rd Qu.:	0.1581		
Max.	:467.7814	Max.	:2120.4743	Max.	: 6.1480	Max.	: 7.6594		

Suppose that we are interested in the association between allergen levels in the mattress and in the living room, under a linear regression model with log transformation in both variables, adjusting for cat ownership. The model can be fitted as follows:

```
> modcat <- tlm (y = logroom, x = logmattress, z = cat, data = feld1, ypow = 0, xpow = 0)
```

where `ypow = 0` and `xpow = 0` indicate that both the explanatory and the response variables are already log transformed. The fitted model provides the following results:

```
> summary(modcat)
```

Linear regression fitted model in the transformed space

-----

Transformations:

```

In the response variable: log
In the explanatory variable: log

```

Call:

```
lm(formula = logroom ~ logmattress + cat, data = feld1)
```

Residuals:

Min	1Q	Median	3Q	Max
-5.9172	-1.1228	-0.0631	0.9464	6.0440

Coefficients:

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	-0.05412	0.11651	-0.465	0.642
logmattress	0.63936	0.04454	14.354	< 2e-16 ***
catyes	1.52747	0.23098	6.613	1.03e-10 ***

---

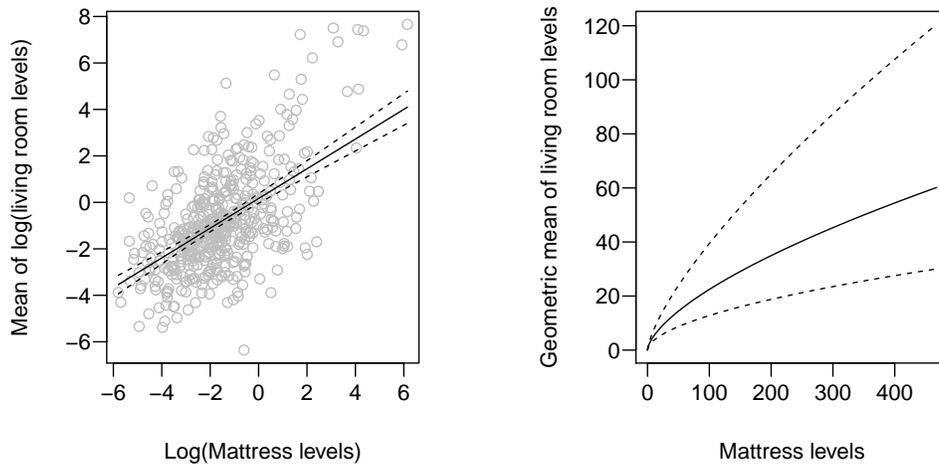
Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 1.68 on 468 degrees of freedom  
 Multiple R-squared: 0.4192, Adjusted R-squared: 0.4167  
 F-statistic: 168.9 on 2 and 468 DF, p-value: < 2.2e-16

We can create Figure 4 with

```
> plot(modcat, type = "transformed", observed = T, xname = "Mattress levels", yname = "living room levels")
> plot(modcat, xname = "Mattress levels", yname = "living room levels")
```

which provides a graphical representation of the relationship between allergen levels in the mattress and in the living room. Measures in Figure 4 have been obtained after averaging the expected measure of  $Y$  over all subjects in the dataset. This is equivalent to saying that measures and effects are calculated for an average individual in the population and they can be interpreted as adjusted measures and adjusted effects [6].



**Figure 4.** Mean of birth weight as a function of cotinine in log scale (left) and in the natural scale (right). Dashed lines represent 95% confidence intervals for the measure.

The function `effectInfo` provides information on interpreting the relationship between allergen levels in mattress ( $X$ ) and in living room ( $Y$ ):

```
> effectInfo(modcat)
```

The effect of  $X$  on  $Y$  can be summarized with a single number as follows:

- Change in  $X$ : multiplicative of factor  $q$  (equivalently, adding an  $r = 100 * (q - 1)\%$  to  $X$ )
- Type of effect on  $Y$ : percent change in the geometric mean of  $Y$
- Effect size:  $100 * (q^{\beta} - 1)\%$

```

beta coefficient estimate:
      Estimate Std. Error t value Pr(>|t|)
logmattress 0.6393565 0.04454319 14.35363 5.610118e-39

```

Further details can be obtained using `effect()`.

In this case, if we use multiplicative (or percent) changes in both variables, a summary effect can be obtained, which is independent of the value of the explanatory variable for which the effect is computed.

The function `effect` provides as default the expected change in allergen levels in the living room for a percent change in allergen levels in mattress equal to the interquartile ratio:

```
> effect(modcat)
```

```
Computing effects...
```

```
Adjusted percent change in the geometric mean of the response variable
for an 'r'% change in the explanatory variable equivalent to the
interquartile ratio:
```

```

      r Estimate lower95% upper95%
1 775.6541 300.393 231.1347 384.137

```

For further information on interpreting the effect use `effectInfo()`.

If we are interested in the effect of cat ownership on allergen levels in the living room, with log transformation in this, we should run a new model with cat ownership as the explanatory variable:

```
> modcat2 <- tlm (y = logroom, x = cat, data = feld1, ypow = 0)
> modcat2
```

```
Linear regression fitted model in the transformed space
```

```
Transformations:
```

```
  In the response variable: log
```

```
Call:
```

```
lm(formula = logroom ~ cat, data = feld1)
```

```
Coefficients:
```

```
(Intercept)      catyes
      -1.218         2.528
```

Then, we can computed measures:

```
> MY(modcat2)
```

Estimated adjusted geometric mean of the response variable in the original space:

```
xlevel geomMean(Y) lower95% upper95%
1 no 0.295888 0.2429528 0.360357
2 yes 3.706041 2.2935543 5.988407
```

and the effect:

```
> effect(modcat2)
```

Computing effects...

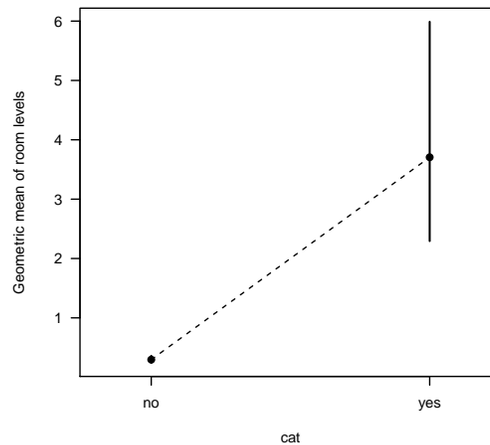
Adjusted change in the geometric mean of the response variable when the explanatory variable changes from its reference level, 'no', to an alternative level (confidence interval for the difference based on 999 bootstrap samples):

```
EstimateDiff lower95% upper95% EstimatePercent lower95% upper95%
no -> yes 3.410153 1.408236 7.556697 1152.515 645.5631 2004.171
```

For further information on interpreting the effect use `effectInfo()`.

A graphical representation of the relationship can be obtained (see Figure 5):

```
> plot(modcat2, yname = "room levels")
```



**Figure 5.** Geometric mean (and 95% confidence intervals) of allergen levels in the living room as a function of cat ownership.

### 3.1.4 Power transformations

Consider now the modeling of the association between triglycerides,  $X$ , and glucose,  $Y$ , levels in blood, both measured in mg/dl. Variables  $X$  and  $Y$  were transformed under power functions  $g(X) = 1/\sqrt{X} = X^{-1/2}$  and  $f(Y) = 1/Y^2 = Y^{-2}$ , respectively, to achieve linearity. The `glucose` data were simulated to emulate true data pattern observed in a real study [2].

First, we can load and explore data (further information about data is available with the help function, `?glucose`):

```
> data(glucose)
> dim(glucose)

[1] 400  4

> head(glucose)

  trigly gluco  inv12tri  inv2glu
1    264   116 0.06154575 7.431629e-05
2    151   123 0.08137885 6.609822e-05
3     67    96 0.12216944 1.085069e-04
4     73    86 0.11704115 1.352082e-04
5    180   104 0.07453560 9.245562e-05
6    130   114 0.08770580 7.694675e-05

> summary(glucose)

      trigly      gluco      inv12tri      inv2glu
Min.   : 33.00   Min.   : 63.00   Min.   :0.03875   Min.   :1.457e-05
1st Qu.: 70.75   1st Qu.: 85.00   1st Qu.:0.08805   1st Qu.:8.573e-05
Median : 94.50   Median : 95.00   Median :0.10287   Median :1.108e-04
Mean   :112.38   Mean   : 99.13   Mean   :0.10297   Mean   :1.131e-04
3rd Qu.:129.00   3rd Qu.:108.00   3rd Qu.:0.11889   3rd Qu.:1.384e-04
Max.   :666.00   Max.   :262.00   Max.   :0.17408   Max.   :2.520e-04
```

Then, we can fit and explore the model of interest:

```
> modglucose <- tlm(y = inv2glu, x = inv12tri, data = glucose, ypow = -2, xpow = -1/2)
> summary(modglucose)
```

Linear regression fitted model in the transformed space

-----  
Transformations:

In the response variable: power, exponent = -2

In the explanatory variable: power, exponent = -1/2

Call:

```
lm(formula = inv2glu ~ inv12tri, data = glucose)
```

Residuals:

Min	1Q	Median	3Q	Max
-9.831e-05	-2.282e-05	-8.110e-07	2.037e-05	1.226e-04

Coefficients:

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	5.424e-05	8.409e-06	6.45	3.25e-10 ***
inv12tri	5.715e-04	7.982e-05	7.16	3.92e-12 ***

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 3.557e-05 on 398 degrees of freedom  
Multiple R-squared: 0.1141, Adjusted R-squared: 0.1119  
F-statistic: 51.27 on 1 and 398 DF, p-value: 3.915e-12

The function `MY` provides a numerical representation of the relationship between triglycerides ( $X$ ) and glucose ( $Y$ ), under the fitted model, in the original scale of the variables:

```
> MY(modglucose)
```

Estimated adjusted median of the response variable in the original space:

	X	median(Y)	lower95%	upper95%
1	33.0000	80.65215	77.74929	83.90648
2	103.3333	95.14538	93.64405	96.72131
3	173.6667	101.21681	98.47839	104.19717
4	244.0000	104.92701	101.08180	109.24731
5	314.3333	107.53510	102.82668	112.95582
6	384.6667	109.51322	104.11239	115.85237
7	455.0000	111.08770	105.11481	118.21011
8	525.3333	112.38364	105.92678	120.18625
9	595.6667	113.47702	106.60300	121.87907
10	666.0000	114.41723	107.17821	123.35386

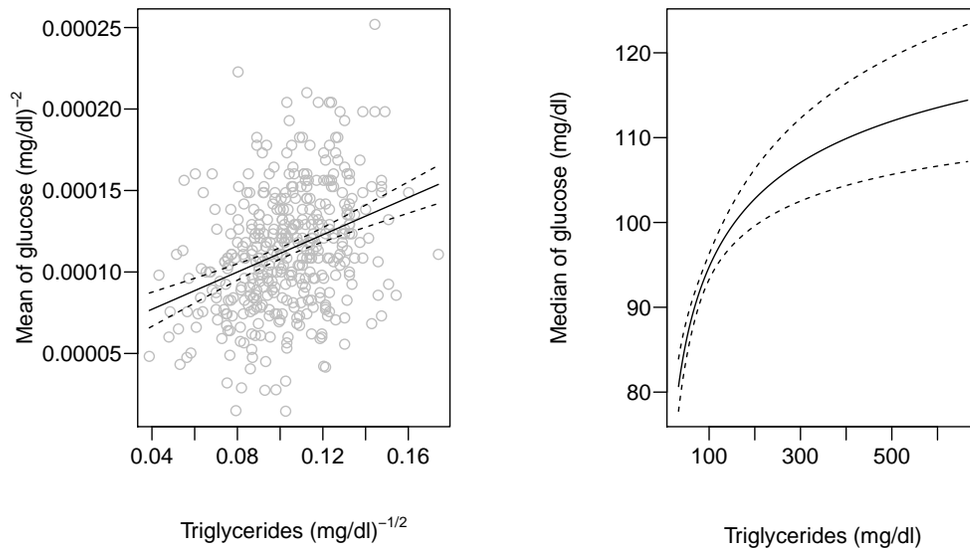
This relationship can also be represented graphically in Figure 6:

```
> plot(modglucose, type = "transformed", observed = T, xname = "Triglycerides (mg/dl)", yname = "glucose (mg/dl)")  
> plot(modglucose, xname = "Triglycerides (mg/dl)", yname = "glucose (mg/dl)")
```

The function `effectInfo` indicates that, under the fitted model, there is no summary effect:

```
> effectInfo(modglucose)
```

The effect of  $X$  on  $Y$  cannot be summarized with a single number. Its behavior can be explored using `effect()`.



**Figure 6.** Relationship between glucose triglycerides levels in the transformed space (left) and in the natural scale of the variables (right). Dashed lines represent 95% confidence intervals for the measure.

Indeed, for general transformations it is not possible to find a summary measure that works for all values of  $X$  and its change. In such cases, we can create tables with the four possible combinations that result when considering both additive and multiplicative changes in both  $X$  and  $Y$ . These tables should report the effects for several basal values of  $X$  along the observed range [7]. For instance, the 2.5th and 97.5th percentiles of triglycerides level were 47 mg/dl and 313 mg/dl, respectively. Thus, we can report the effects between pairs of consecutive values of  $X = 50, 100, 150, 200$  and 250 mg/dl, that is to say, for an additive increase of  $c = 50$  mg/dl, and also between pairs of consecutive values: 50, 75, 112.5, 168.8 and 253.1 mg/dl, that is, for an  $r = 50\%$  increase:

```
> # Effects for an additive change in triglycerides level:
> xc <- 50 * (1:5)
> xc
```

```
[1] 50 100 150 200 250
```

```
> effectXdif <- effect(modglucose, x1 = xc)
```

Computing effects...

```
> effectXdif
```

Adjusted change in the median of the response variable when the explanatory variable changes from  $x_1$  to  $x_2$  (confidence intervals based on 999 bootstrap

samples):

	x1	x2	EstimateDiff	lower95%	upper95%	EstimatePercent	lower95%	upper95%
1	50	100	8.703222	6.334019	10.907257	10.114803	7.250154	12.960938
2	100	150	4.802290	3.313412	6.464551	5.068507	3.506924	6.792880
3	150	200	3.235200	2.178028	4.439836	3.249826	2.215835	4.370665
4	200	250	2.397980	1.602089	3.372809	2.333003	1.597478	3.172078

> # Effects for an percent change in triglycerides level:

> xq <- 50 \* 1.5^(0:4)

> xq

[1] 50.000 75.000 112.500 168.750 253.125

> effectXperc <- effect(modglucose, x1 = xq)

Computing effects...

> effectXperc

Adjusted change in the median of the response variable when the explanatory variable changes from x1 to x2 (confidence intervals based on 999 bootstrap samples):

	x1	x2	EstimateDiff	lower95%	upper95%	EstimatePercent	lower95%	upper95%
1	50.00	75.000	5.152950	3.813373	6.363338	5.988710	4.361787	7.553217
2	75.00	112.500	4.971764	3.542537	6.461374	5.451653	3.853272	7.140190
3	112.50	168.750	4.724024	3.260444	6.322945	4.912204	3.396935	6.507858
4	168.75	253.125	4.420430	2.976703	6.163127	4.381299	3.006967	5.941818

Exploring previous results, one can see that the more easily interpretable effect appears to be the additive change in the median of glucose level associated to a percent change in triglycerides level. Indeed, we can see that, for any given value of the triglycerides level along the observed range, a 50% increase in triglycerides level is associated to around a 4.8 mg/dl increase in the median glucose level. For the other measures, the effect is more dependent on the basal value of the triglycerides level.

### 3.2 Logistic regression model with log transformation in the explanatory variable

Revisiting the cotinine example -section 3.1.2-, suppose we are now interested in the association between low birth weight -defined as weight lower than 2500 g-,  $Y$ , and cotinine level,  $X$ , after log transforming  $X$ . The model can be fitted as follows:

```
> modcot2 <- tlm(y = underweight, x = logcotinine, data = cotinine, xpow = 0, family = binomial)
```

where  $xpow = 0$  indicates that the explanatory variable is already log transformed and the argument `family = binomial` indicates that the regression model is logistic with logit link (default is `family = gaussian`, for the lineal regression model). The fitted model provides the following results:

```
> summary(modcot2)
```

Logistic regression fitted model in the transformed space

-----  
Transformations:

In the response variable: logit link for logistic regression

In the explanatory variable: log

Call:

```
glm(formula = underweight ~ logcotinine, family = binomial, data = cotinine)
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-0.7061	-0.4186	-0.3231	-0.2821	2.6239

Coefficients:

	Estimate	Std. Error	z value	Pr(> z )
(Intercept)	-3.5146	0.4539	-7.744	9.65e-15 ***
logcotinine	0.3306	0.1289	2.566	0.0103 *

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 175.09 on 350 degrees of freedom

Residual deviance: 168.45 on 349 degrees of freedom

AIC: 172.45

Number of Fisher Scoring iterations: 5

Then, we can obtain the probability of low birth weight as a function of cotinine level:

```
> MY(modcot2)
```

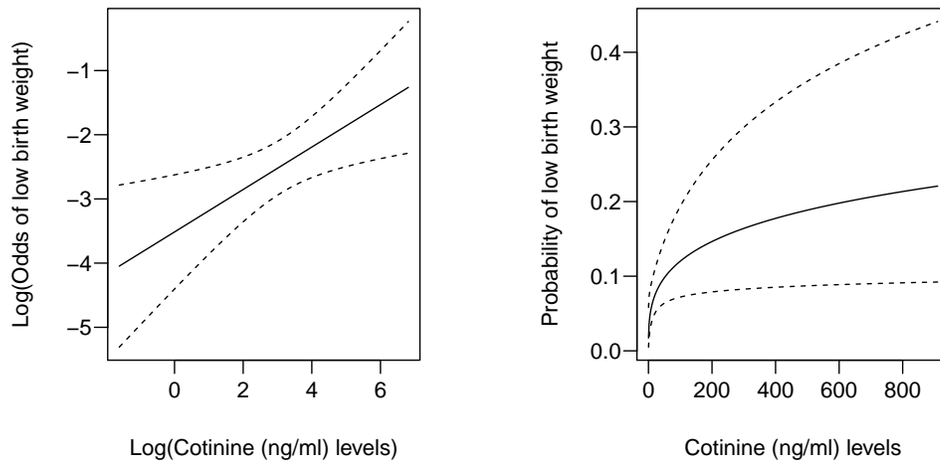
Estimated adjusted probability of the response variable in the original space:

	X	P(Y)	lower95%	upper95%
1	0.2000	0.01717866	0.004920113	0.05819342
2	101.2889	0.12049711	0.072138318	0.19447904
3	202.3778	0.14693118	0.079055696	0.25683106
4	303.4667	0.16452717	0.082757167	0.30061256
5	404.5556	0.17801453	0.085288433	0.33466844
6	505.6444	0.18906298	0.087213645	0.36260410
7	606.7333	0.19847559	0.088768165	0.38629112
8	707.8222	0.20670597	0.090072508	0.40684022
9	808.9111	0.21403755	0.091196603	0.42496992
10	910.0000	0.22066025	0.092184631	0.44117414

A graphical representation of the relationship between cotinine level and low birth weight can be obtained by:

```
> plot(modcot2, type = "transformed", xname = "Cotinine (ng/ml) levels", yname = "low birth weigh
> plot(modcot2, xname = "Cotinine (ng/ml) levels", yname = "low birth weight")
```

which provides Figure 7.



**Figure 7.** Log odds of low birth weight as a function of log cotinine (left) and probability of low birth weight as a function of cotinine level (right). Dashed lines represent 95% confidence intervals for the measure.

Regarding effects, the function `effectInfo` indicates that, under the fitted model, we can summarize the effect of cotinine level ( $X$ ) on low birth weight ( $Y$ ) in terms of odds ratio (OR) for a percent (or multiplicative) change in cotinine level:

```
> effectInfo(modcot2)
```

The effect of  $X$  on  $Y$  can be summarized with a single number as follows:

- Change in  $X$ : multiplicative of factor  $q$  (equivalently, adding an  $r = 100 * (q - 1)\%$  to  $X$ )
- Type of effect on  $Y$ : odds ratio of  $Y$
- Effect size:  $q^{\beta}$

beta coefficient estimate:

	Estimate	Std. Error	z value	Pr(> z )
logcotinine	0.3306394	0.1288549	2.565982	0.01028842

Further details can be obtained using `effect()`.

Thus, the function `effect` provides as default the OR of low birth weight for a percent change in the cotinine level equal to the interquartile ratio:

```
> effect(modcot2)
```

Computing effects...

Adjusted odds ratio of the response variable for an 'r'% change in the explanatory variable equivalent to the interquartile ratio:

```
      r Estimate lower95% upper95%
1 1112.878 2.282194 1.21516 4.286193
```

For further information on interpreting the effect use `effectInfo()`.

Alternatively, the effect for a 10-fold change in the cotinine level is:

```
> effect(modcot2, q = 10)
```

Computing effects...

Adjusted odds ratio of the response variable for an 'r' = 900% change in the explanatory variable:

```
      r Estimate lower95% upper95%
1 900 2.141112 1.196984 3.829926
```

For further information on interpreting the effect use `effectInfo()`.

## References

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