Testing covariate modelling in hierarchical parent degradation kinetics with residue data on mesotrione

Johannes Ranke

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Introduction

The purpose of this document is to test demonstrate how nonlinear hierarchical models (NLHM) based on the parent degradation models SFO, FOMC, DFOP and HS can be fitted with the mkin package, also considering the influence of covariates like soil pH on different degradation parameters. Because in some other case studies, the SFORB parameterisation of biexponential decline has shown some advantages over the DFOP parameterisation, SFORB was included in the list of tested models as well.

The mkin package is used in version 1.2.4, which is contains the functions that were used for the evaluations. The saemix package is used as a backend for fitting the NLHM, but is also loaded to make the convergence plot function available.

This document is processed with the knitr package, which also provides the kable function that is used to improve the display of tabular data in R markdown documents. For parallel processing, the parallel package is used.

```
library(mkin)
library(saemix)
library(parallel)
n_cores <- detectCores()
if (Sys.info()["sysname"] == "Windows") {
   cl <- makePSOCKcluster(n_cores)
} else {
   cl <- makeForkCluster(n_cores)
}</pre>
```

Test data

```
data_file <- system.file(
  "testdata", "mesotrione_soil_efsa_2016.xlsx", package = "mkin")
meso_ds <- read_spreadsheet(data_file, parent_only = TRUE)</pre>
```

The following tables show the covariate data and the 18 datasets that were read in from the spreadsheet file.

```
pH <- attr(meso_ds, "covariates")
kable(pH, caption = "Covariate data")</pre>
```

Table 1: Covariate data

	рН
Richmond	6.2
Richmond 2	6.2
ERTC	6.4
Toulouse	7.7
Picket Piece	7.1
721	5.6
722	5.7
723	5.4
724	4.8
725	5.8
727	5.1
728	5.9
729	5.6
730	5.3
731	6.1
732	5.0
741	5.7
742	7.2

```
for (ds_name in names(meso_ds)) {
  print(
    kable(mkin_long_to_wide(meso_ds[[ds_name]]),
        caption = paste("Dataset", ds_name),
        booktabs = TRUE, row.names = FALSE))
}
```

Table 2: Dataset Richmond

time	meso
0.000000	91.00
1.179050	86.70
3.537149	73.60
7.074299	61.50
10.611448	55.70
15.327647	47.70
17.685747	39.50
24.760046	29.80
35.371494	19.60
68.384889	5.67
0.000000	97.90
1.179050	96.40
3.537149	89.10
7.074299	74.40
10.611448	57.40
15.327647	46.30
18.864797	35.50
27.118146	27.20
35.371494	19.10
74.280138	6.50
108.472582	3.40
142.665027	2.20

Table 3: Dataset Richmond 2

time	meso
0.000000	96.0
2.422004	82.4
5.651343	71.2
8.073348	53.1
11.302687	48.5
16.954030	33.4
22.605373	24.2
45.210746	11.9

Table 4: Dataset ERTC

time	meso
0.000000	99.9
2.755193	80.0
6.428782	42.1
9.183975	50.1

time	meso
12.857565	28.4
19.286347	39.8
25.715130	29.9
51.430259	2.5

Table 5: Dataset Toulouse

time	meso
0.000000	96.8
2.897983	63.3
6.761960	22.3
9.659942	16.6
13.523919	16.1
20.285879	17.2
27.047838	1.8

Table 6: Dataset Picket Piece

time	meso
0.000000	102.0
2.841195	73.7
6.629454	35.5
9.470649	31.8
13.258909	18.0
19.888364	3.7

Table 7: Dataset 721

	time	meso
	0.00000	86.4
	11.24366	61.4
-	22.48733	49.8
,	33.73099	41.0
4	44.97466	35.1
-		

Table 8: Dataset 722

$_{ m time}$	meso
0.00000	90.3
11.24366	52.1
22.48733	37.4
33.73099	21.2
44.97466	14.3

Table 9: Dataset 723

time	meso
0.00000	89.3
11.24366	70.8
22.48733	51.1
33.73099	42.7
44.97466	26.7

Table 10: Dataset 724

time	meso
0.000000	89.4
9.008208	65.2
18.016415	55.8
27.024623	46.0
36.032831	41.7

Table 11: Dataset 725

time	meso
0.00000	89.0
10.99058	35.4
21.98116	18.6
32.97174	11.6
43.96232	7.6

Table 12: Dataset 727

time	meso
0.00000	91.3
10.96104	63.2
21.92209	51.1
32.88313	42.0
43.84417	40.8

Table 13: Dataset 728

time	mesc
0.00000	91.8
11.24366	43.6
22.48733	22.0
33.73099	15.9
44.97466	8.8

Table 14: Dataset 729

time	meso
0.00000	91.6
11.24366	60.5
22.48733	43.5
33.73099	28.4
44.97466	20.5
44.97466	20.5

Table 15: Dataset 730

time	meso
0.00000	92.7
11.07446	58.9
22.14893	44.0
33.22339	46.0
44.29785	29.3

Table 16: Dataset 731

time	meso
0.00000	92.1
11.24366	64.4
22.48733	45.3
33.73099	33.6
44.97466	23.5
11.24366 22.48733 33.73099	64.4 45.3 33.6

Table 17: Dataset 732

time	meso
0.00000	90.3
11.24366	58.2
22.48733	40.1
33.73099	33.1
44.97466	25.8

Table 18: Dataset 741

time	meso
0.00000	90.3
10.84712	68.7
21.69424	58.0
32.54136	52.2
43.38848	48.0

Table 19: Dataset 742

time	meso
0.00000	92.0
11.24366	60.9
22.48733	36.2
33.73099	18.3
44.97466	8.7

Separate evaluations

In order to obtain suitable starting parameters for the NLHM fits, separate fits of the five models to the data for each soil are generated using the mmkin function from the mkin package. In a first step, constant variance is assumed. Convergence is checked with the status function.

```
deg_mods <- c("SFO", "FOMC", "DFOP", "SFORB", "HS")
f_sep_const <- mmkin(
  deg_mods,
  meso_ds,
  error_model = "const",
  cluster = cl,
  quiet = TRUE)</pre>
```

status(f_sep_const[, 1:5]) |> kable()

	Richmond	Richmond 2	ERTC	Toulouse	Picket Piece
SFO	OK	OK	OK	OK	OK
FOMC	OK	OK	OK	OK	\mathbf{C}
DFOP	OK	OK	OK	OK	OK
SFORB	OK	OK	OK	OK	OK
HS	OK	OK	\mathbf{C}	OK	OK

status(f_sep_const[, 6:18]) |> kable()

	721	722	723	724	725	727	728	729	730	731	732	741	742
SFO	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK
FOMC	OK	OK	\mathbf{C}	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK
DFOP	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK
SFORB	OK	OK	OK	OK	OK	OK	OK	\mathbf{C}	OK	OK	OK	OK	OK
HS	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK

In the tables above, OK indicates convergence and C indicates failure to converge. Most separate fits with constant variance converged, with the exception of two FOMC fits, one SFORB fit and one HS fit.

```
f_sep_tc <- update(f_sep_const, error_model = "tc")
status(f_sep_tc[, 1:5]) |> kable()
```

	Richmond	Richmond 2	ERTC	Toulouse	Picket Piece
SFO	ОК	ОК	OK	OK	OK
FOMC	OK	OK	OK	OK	OK
DFOP	\mathbf{C}	OK	OK	OK	OK
SFORB	OK	OK	OK	OK	OK
HS	OK	OK	\mathbf{C}	OK	OK

```
status(f_sep_tc[, 6:18]) |> kable()
```

	721	722	723	724	725	727	728	729	730	731	732	741	742
SFO	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK	OK
FOMC	OK	OK	\mathbf{C}	OK	\mathbf{C}	\mathbf{C}	OK	\mathbf{C}	OK	\mathbf{C}	OK	\mathbf{C}	OK
DFOP	\mathbf{C}	OK	OK	OK	\mathbf{C}	OK	OK	OK	OK	\mathbf{C}	OK	\mathbf{C}	OK

	721	722	723	724	725	727	728	729	730	731	732	741	742
SFORB	C	OK	OK	OK	C	OK	OK	C	OK	OK	OK	C	OK
HS	OK	C	OK	OK	OK								

With the two-component error model, the set of fits that did not converge is larger, with convergence problems appearing for a number of non-SFO fits.

Hierarchical model fits without covariate effect

The following code fits hierarchical kinetic models for the ten combinations of the five different degradation models with the two different error models in parallel.

```
f_saem_1 <- mhmkin(list(f_sep_const, f_sep_tc), cluster = cl)
status(f_saem_1) |> kable()
```

	const	tc
SFO	OK	OK
FOMC	OK	OK
DFOP	OK	OK
SFORB	OK	OK
HS	OK	OK

All fits terminate without errors (status OK).

anova(f_saem_1) |> kable(digits = 1)

	npar	AIC	BIC	Lik
SFO const	5	800.0	804.5	-395.0
SFO tc	6	801.9	807.2	-394.9
FOMC const	7	787.4	793.6	-386.7
FOMC tc	8	788.9	796.1	-386.5
DFOP const	9	787.6	795.6	-384.8
SFORB const	9	787.4	795.4	-384.7
HS const	9	781.9	789.9	-382.0
DFOP tc	10	787.4	796.3	-383.7
SFORB tc	10	795.8	804.7	-387.9
HS tc	10	783.7	792.7	-381.9

The model comparisons show that the fits with constant variance are consistently preferable to the corresponding fits with two-component error for these data. This is confirmed by the fact that the parameter b.1 (the relative standard deviation in the fits obtained with the saemix package), is ill-defined in all fits.

illparms(f_saem_1) |> kable()

	const	tc
SFO	$sd(meso_0)$	$sd(meso_0), b.1$
FOMC	$sd(meso_0), sd(log_beta)$	sd(meso_0), sd(log_beta), b.1
DFOP	$sd(meso_0), sd(log_k1)$	$sd(meso_0), sd(g_qlogis), b.1$
SFORB	sd(meso_free_0), sd(log_k_meso_free_bound)	sd(meso_free_0), sd(log_k_meso_free_bound), b.1
HS	$sd(meso_0)$	$sd(meso_0), b.1$

For obtaining fits with only well-defined random effects, we update the set of fits, excluding random effects that were ill-defined according to the <code>illparms</code> function.

```
f_saem_2 <- update(f_saem_1, no_random_effect = illparms(f_saem_1))
status(f_saem_2) |> kable()
```

	const	tc
SFO	OK	OK
FOMC	OK	OK

	const	tc
DFOP	OK	OK
SFORB	OK	OK
HS	OK	OK

The updated fits terminate without errors.

illparms(f_saem_2) |> kable()

	const	tc
SFO		b.1
FOMC		b.1
DFOP		b.1
SFORB		b.1
HS		

No ill-defined errors remain in the fits with constant variance.

Hierarchical model fits with covariate effect

In the following sections, hierarchical fits including a model for the influence of pH on selected degradation parameters are shown for all parent models. Constant variance is selected as the error model based on the fits without covariate effects. Random effects that were ill-defined in the fits without pH influence are excluded. A potential influence of the soil pH is only included for parameters with a well-defined random effect, because experience has shown that for such parameters no significant pH effect could be found.

SFO

```
sfo_pH <- saem(f_sep_const["SFO", ], no_random_effect = "meso_0", covariates = pH,
    covariate_models = list(log_k_meso ~ pH))
summary(sfo_pH)$confint_trans |> kable(digits = 2)
```

	est.	lower	upper
meso_0	91.35	89.27	93.43
log_k_meso	-6.66	-7.97	-5.35
beta_pH(log_k_meso)	0.59	0.37	0.81
a.1	5.48	4.71	6.24
$SD.log_k_meso$	0.35	0.23	0.47

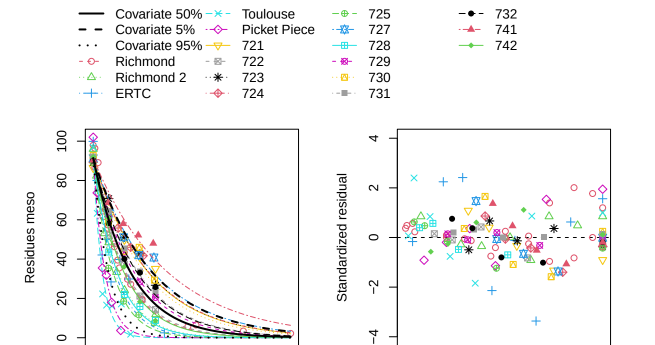
The parameter showing the pH influence in the above table is beta_pH(log_k_meso). Its confidence interval does not include zero, indicating that the influence of soil pH on the log of the degradation rate constant is significantly greater than zero.

```
anova(f_saem_2[["SFO", "const"]], sfo_pH, test = TRUE)
```

Data: 116 observations of 1 variable(s) grouped in 18 datasets

The comparison with the SFO fit without covariate effect confirms that considering the soil pH improves the model, both by comparison of AIC and BIC and by the likelihood ratio test.

plot(sfo_pH)



Endpoints for a model with covariates are by default calculated for the median of the covariate values. This quantile can be adapted, or a specific covariate value can be given as shown below.

0

20

40

Predicted

60

80

120

endpoints(sfo_pH)

20

40

60 80

Time

```
$covariates
      рΗ
50% 5.75
$distimes
         DT50
                  DT90
meso 18.52069 61.52441
endpoints(sfo_pH, covariate_quantile = 0.9)
$covariates
      рΗ
90% 7.13
$distimes
         DT50
                  DT90
meso 8.237019 27.36278
endpoints(sfo_pH, covariates = c(pH = 7.0))
$covariates
     рΗ
User 7
$distimes
        DT50
                DT90
```

FOMC

```
fomc_pH <- saem(f_sep_const["FOMC", ], no_random_effect = "meso_0", covariates = pH,
    covariate_models = list(log_alpha ~ pH))</pre>
```

summary(fomc_pH)\$confint_trans |> kable(digits = 2)

	est.	lower	upper
meso_0	92.84	90.75	94.93
log_alpha	-2.21	-3.49	-0.92
$beta_pH(log_alpha)$	0.58	0.37	0.79
log_beta	4.21	3.44	4.99
a.1	5.03	4.32	5.73
SD.log_alpha	0.00	-23.77	23.78
SD.log_beta	0.37	0.01	0.74

As in the case of SFO, the confidence interval of the slope parameter (here beta_pH(log_alpha)) quantifying the influence of soil pH does not include zero, and the model comparison clearly indicates that the model with covariate influence is preferable. However, the random effect for alpha is not well-defined any more after inclusion of the covariate effect (the confidence interval of SD.log_alpha includes zero).

```
illparms(fomc_pH)
```

[1] "sd(log_alpha)"

Therefore, the model is updated without this random effect, and no ill-defined parameters remain.

```
fomc_pH_2 <- update(fomc_pH, no_random_effect = c("meso_0", "log_alpha"))
illparms(fomc_pH_2)</pre>
```

```
anova(f_saem_2[["FOMC", "const"]], fomc_pH, fomc_pH_2, test = TRUE)
```

Data: 116 observations of 1 variable(s) grouped in 18 datasets

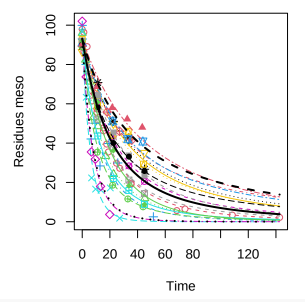
Model comparison indicates that including pH dependence significantly improves the fit, and that the reduced model with covariate influence results in the most preferable FOMC fit.

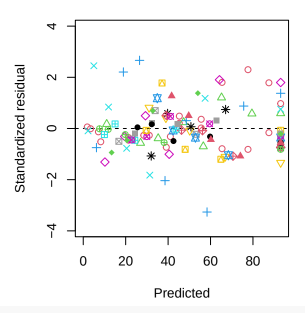
```
summary(fomc_pH_2)$confint_trans |> kable(digits = 2)
```

	est.	lower	upper
meso_0	93.05	90.98	95.13
log_alpha	-2.91	-4.18	-1.63
$beta_pH(log_alpha)$	0.66	0.44	0.87
log_beta	3.95	3.29	4.62
a.1	4.98	4.28	5.68
$SD.log_beta$	0.40	0.26	0.54

plot(fomc_pH_2)







endpoints(fomc_pH_2)

\$covariates

рΗ

50% 5.75

\$distimes

DT50 DT90 DT50back meso 17.30248 82.91343 24.95943

endpoints(fomc_pH_2, covariates = c(pH = 7))

\$covariates

рΗ

User 7

\$distimes

DT50 DT90 DT50back meso 6.986239 27.02927 8.136621

DFOP

In the DFOP fits without covariate effects, random effects for two degradation parameters (k2 and g) were identifiable. summary(f_saem_2[["DFOP", "const"]])\$confint_trans |> kable(digits = 2)

	est.	lower	upper
meso_0	93.61	91.58	95.63
log_k1	-1.53	-2.27	-0.79
\log_k2	-3.42	-3.73	-3.11
g_qlogis	-1.67	-2.57	-0.77
a.1	4.74	4.02	5.45
$SD.log_k2$	0.60	0.38	0.81
$SD.g_qlogis$	0.94	0.33	1.54

A fit with pH dependent degradation parameters was obtained by excluding the same random effects as in the refined DFOP fit without covariate influence, and including covariate models for the two identifiable parameters k2 and g.

```
dfop_pH <- saem(f_sep_const["DFOP", ], no_random_effect = c("meso_0", "log_k1"),
    covariates = pH,
    covariate_models = list(log_k2 ~ pH, g_qlogis ~ pH))</pre>
```

The corresponding parameters for the influence of soil pH are beta_pH(log_k2) for the influence of soil pH on k2, and beta_pH(g_qlogis) for its influence on g.

summary(dfop_pH)\$confint_trans |> kable(digits = 2)

	est.	lower	upper
meso_0	92.84	90.85	94.84
log_k1	-2.82	-3.09	-2.54
\log_k2	-11.48	-15.32	-7.64
$beta_pH(log_k2)$	1.31	0.69	1.92
g_qlogis	3.13	0.47	5.80
$beta_pH(g_qlogis)$	-0.57	-1.04	-0.09
a.1	4.96	4.26	5.65
SD.log_k2	0.76	0.47	1.05
SD.g_qlogis	0.01	-9.96	9.97

illparms(dfop_pH)

[1] "sd(g_qlogis)"

Confidence intervals for neither of them include zero, indicating a significant difference from zero. However, the random effect for ${\sf g}$ is now ill-defined. The fit is updated without this ill-defined random effect.

```
dfop_pH_2 <- update(dfop_pH,
    no_random_effect = c("meso_0", "log_k1", "g_qlogis"))
illparms(dfop_pH_2)</pre>
```

[1] "beta_pH(g_qlogis)"

Now, the slope parameter for the pH effect on g is ill-defined. Therefore, another attempt is made without the corresponding covariate model.

```
dfop_pH_3 <- saem(f_sep_const["DFOP", ], no_random_effect = c("meso_0", "log_k1"),
    covariates = pH,
    covariate_models = list(log_k2 ~ pH))
illparms(dfop_pH_3)</pre>
```

```
[1] "sd(g_qlogis)"
```

As the random effect for g is again ill-defined, the fit is repeated without it.

```
dfop_pH_4 <- update(dfop_pH_3, no_random_effect = c("meso_0", "log_k1", "g_qlogis"))
illparms(dfop_pH_4)</pre>
```

While no ill-defined parameters remain, model comparison suggests that the previous model dfop_pH_2 with two pH dependent parameters is preferable, based on information criteria as well as based on the likelihood ratio test.

```
anova(f_saem_2[["DFOP", "const"]], dfop_pH_2, dfop_pH_3, dfop_pH_4)
```

Data: 116 observations of 1 variable(s) grouped in 18 datasets

```
npar AIC BIC Lik
f_saem_2[["DFOP", "const"]] 7 782.94 789.18 -384.47
dfop_pH_4 7 767.35 773.58 -376.68
dfop_pH_2 8 765.14 772.26 -374.57
dfop_pH_3 8 769.00 776.12 -376.50
dfop_pH 9 769.10 777.11 -375.55
anova(dfop_pH_2, dfop_pH_4, test = TRUE)
```

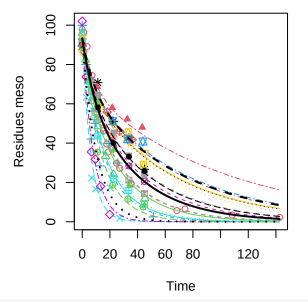
Data: 116 observations of 1 variable(s) grouped in 18 datasets

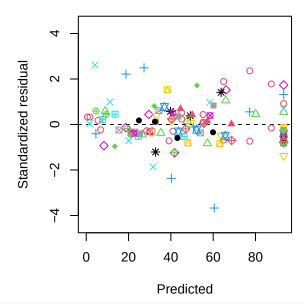
```
npar AIC BIC Lik Chisq Df Pr(>Chisq)
dfop_pH_4 7 767.35 773.58 -376.68
dfop_pH_2 8 765.14 772.26 -374.57 4.2153 1 0.04006 *
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Here, the reduced model <code>dfop_pH_4</code> is still proposed to be preferable model, as all parameters are clearly identifiable, and it will be more easily useable because of the lower complexity.

plot(dfop_pH_4)







endpoints(dfop_pH_4)

\$covariates

рΗ

50% 5.75

\$distimes

endpoints(dfop_pH_4, covariates = c(pH = 7))

\$covariates

рΗ

User 7

\$distimes

SFORB

```
sforb_pH <- saem(f_sep_const["SFORB", ], no_random_effect = c("meso_free_0", "log_k_meso_free_bound"),
    covariates = pH,
    covariate_models = list(log_k_meso_free ~ pH, log_k_meso_bound_free ~ pH))
summary(sforb_pH)$confint_trans |> kable(digits = 2)
```

	est.	lower	upper
meso_free_0	93.42	91.32	95.52
log_k_meso_free	-5.37	-6.94	-3.81
beta_pH(log_k_meso_free)	0.42	0.18	0.67
log_k_meso_free_bound	-3.49	-4.92	-2.05
log_k_meso_bound_free	-9.98	-19.22	-0.74
beta_pH(log_k_meso_bound_free)	1.23	-0.21	2.67
a.1	4.90	4.18	5.63
SD.log_k_meso_free	0.35	0.23	0.47
$SD.log_k_meso_bound_free$	0.13	-1.95	2.20

The confidence interval of beta_pH(log_k_meso_bound_free) includes zero, indicating that the influence of soil pH on k_meso_bound_free cannot reliably be quantified. Also, the confidence interval for the random effect on this parameter (SD.log_k_meso_bound_free) includes zero.

Using the illparms function, these ill-defined parameters can be found more conveniently.

illparms(sforb_pH)

```
[1] "sd(log_k_meso_bound_free)" "beta_pH(log_k_meso_bound_free)"
```

To remove the ill-defined parameters, a second variant of the SFORB model with pH influence is fitted. No ill-defined parameters remain.

```
sforb_pH_2 <- update(sforb_pH,
    no_random_effect = c("meso_free_0", "log_k_meso_free_bound", "log_k_meso_bound_free"),
    covariate_models = list(log_k_meso_free ~ pH))
illparms(sforb_pH_2)</pre>
```

The model comparison of the SFORB fits includes the refined model without covariate effect, and both versions of the SFORB fit with covariate effect.

```
anova(f_saem_2[["SFORB", "const"]], sforb_pH, sforb_pH_2, test = TRUE)
```

Data: 116 observations of 1 variable(s) grouped in 18 datasets

```
npar AIC BIC Lik Chisq Df Pr(>Chisq)
f_saem_2[["SFORB", "const"]] 7 783.40 789.63 -384.70
sforb_pH_2 7 770.94 777.17 -378.47 12.4616 0
sforb_pH 9 768.81 776.83 -375.41 6.1258 2 0.04675 *
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The first model including pH influence is preferable based on information criteria and the likelihood ratio test. However, as it is not fully identifiable, the second model is selected.

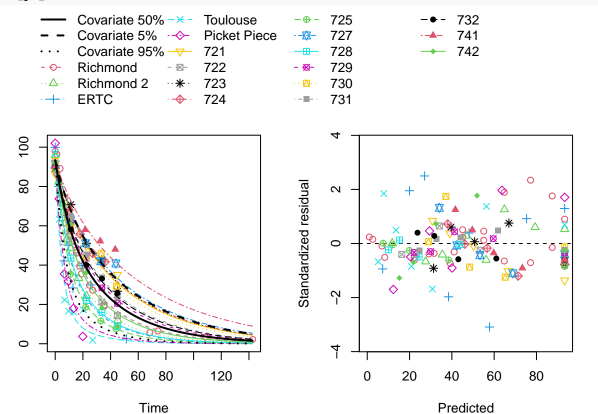
```
summary(sforb_pH_2)$confint_trans |> kable(digits = 2)
```

			est.	lower	upper
meso	free	0	93.32	91.16	95.48

	est.	lower	upper
log_k_meso_free	-6.15	-7.43	-4.86
beta_pH(log_k_meso_free)	0.54	0.33	0.75
log_k_meso_free_bound	-3.80	-5.20	-2.40
$\log_k_{meso_bound_free}$	-2.95	-4.26	-1.64
a.1	5.08	4.38	5.79
SD.log_k_meso_free	0.33	0.22	0.45

```
plot(sforb_pH_2)
```

Residues meso



endpoints(sforb_pH_2)

```
$covariates
      рΗ
50% 5.75
$ff
meso_free
$SFORB
   meso_b1
              meso_b2
                          meso_g
0.09735824 \ 0.02631699 \ 0.31602120
$distimes
         DT50
                  DT90 DT50back DT50_meso_b1 DT50_meso_b2
meso 16.86549 73.15824 22.02282
                                     7.119554
                                                   26.33839
endpoints(sforb_pH_2, covariates = c(pH = 7))
```

```
$covariates
```

pH User 7 \$ff meso_free 1

\$SFORB

meso_b1 meso_b2 meso_g

0.13315233 0.03795988 0.61186191

\$distimes

DT50 DT90 DT50back DT50_meso_b1 DT50_meso_b2 meso 7.932495 36.93311 11.11797 5.205671 18.26

HS

```
hs_pH <- saem(f_sep_const["HS", ], no_random_effect = c("meso_0"),
    covariates = pH,
    covariate_models = list(log_k1 ~ pH, log_k2 ~ pH, log_tb ~ pH))</pre>
```

illparms(hs_pH)

```
[1] "sd(log_tb)" "beta_pH(log_tb)"
```

According to the output of the illparms function, the influence of soil pH on the break time tb cannot reliably be quantified. Thus, the fit is repeated without the corresponding covariate model.

```
hs_pH_2 <- update(hs_pH, covariate_models = list(log_k1 ~ pH, log_k2 ~ pH))
illparms(hs_pH_2)</pre>
```

Model comparison confirms that this model is preferable to the fit without covariate influence, and also to the first version with covariate influence.

```
anova(f_saem_2[["HS", "const"]], hs_pH, hs_pH_2, test = TRUE)
```

Data: 116 observations of 1 variable(s) grouped in 18 datasets

```
npar AIC BIC Lik Chisq Df Pr(>Chisq)

f_saem_2[["HS", "const"]] 8 780.08 787.20 -382.04

hs_pH_2 10 766.47 775.37 -373.23 17.606 2 0.0001503 ***

hs_pH 11 769.80 779.59 -373.90 0.000 1 1.0000000

---

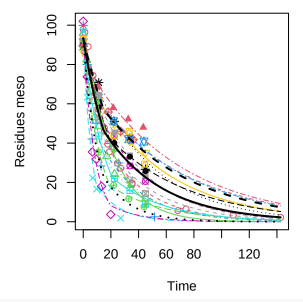
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

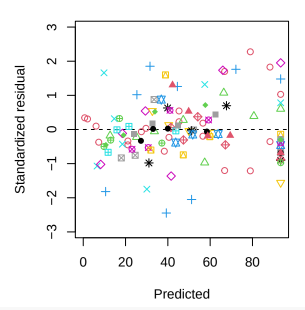
summary(hs_pH_2)$confint_trans |> kable(digits = 2)
```

	est.	lower	upper
meso_0	93.33	91.50	95.15
log_k1	-5.68	-7.09	-4.27
$beta_pH(log_k1)$	0.46	0.22	0.69
log_k2	-6.61	-8.34	-4.88
$beta_pH(log_k2)$	0.50	0.21	0.79
\log_{tb}	2.70	2.33	3.08
a.1	4.45	3.74	5.16
SD.log_k1	0.36	0.22	0.49
$SD.log_k2$	0.23	0.02	0.43
$SD.log_tb$	0.55	0.25	0.85

plot(hs_pH_2)







732

741 742

endpoints(hs_pH_2)

\$covariates

рΗ

50% 5.75

\$distimes

endpoints(hs_pH_2, covariates = c(pH = 7))

\$covariates

рΗ

User 7

\$distimes

DT50 DT90 DT50back DT50_k1 DT50_k2 meso 8.298536 38.85371 11.69613 8.298536 15.71561

Comparison across parent models

After model reduction for all models with pH influence, they are compared with each other.

```
anova(sfo_pH, fomc_pH_2, dfop_pH_2, sforb_pH_2, hs_pH_2)
```

Data: 116 observations of 1 variable(s) grouped in 18 datasets

```
        npar
        AIC
        BIC
        Lik

        sfo_pH
        5
        783.09
        787.54
        -386.54

        fomc_pH_2
        6
        767.49
        772.83
        -377.75

        sforb_pH_2
        7
        770.94
        777.17
        -378.47

        dfop_pH_2
        8
        765.14
        772.26
        -374.57

        hs_pH_2
        10
        766.47
        775.37
        -373.23
```

The DFOP model with pH influence on k2 and g and a random effect only on k2 is finally selected as the best fit.

The endpoints resulting from this model are listed below. Please refer to the Appendix for a detailed listing.

```
endpoints(dfop_pH_2)
```

\$covariates

meso 8.346428 28.34437 8.532507 4.191901 8.753618

Conclusions

These evaluations demonstrate that covariate effects can be included for all types of parent degradation models. These models can then be further refined to make them fully identifiable.

Appendix

Hierarchical fit listings

Fits without covariate effects

Listing 1: Hierarchical SFO fit with constant variance

```
saemix version used for fitting: 3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting: 4.3.1
Date of fit: Fri Aug 4 12:14:17 2023
Date of summary: Fri Aug 4 12:23:55 2023
Equations:
d_{meso}/dt = - k_{meso} * meso
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 6.418 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
    meso_0 log_k_meso
    90.832
               -3.192
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
          meso_0 log_k_meso
                      0.0000
meso_0
              6.752
log_k_meso 0.000
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
  AIC BIC logLik
  800 804.5 -395
Optimised parameters:
                  est.
                            lower upper
meso_0 92.0705 89.9917 94.1493

      log_k_meso
      -3.1641
      -3.4286
      -2.8996

      a.1
      5.4628
      4.6421
      6.2835

      SD.meso_0
      0.0611
      -98.3545
      98.4767

SD.log_k_meso 0.5616 0.3734 0.7499
Correlation:
log_k_meso 0.1132
Random effects:
est. lower upper SD.meso_0 0.0611 -98.3545 98.4767
SD.log_k_meso 0.5616 0.3734 0.7499
Variance model:
     est. lower upper
a.1 5.463 4.642 6.284
{\tt Backtransformed\ parameters:}
            est.
                     lower
meso_0 92.07053 89.99172 94.14933
k_meso 0.04225 0.03243 0.05505
Estimated disappearance times:
     DT50 DT90
meso 16.41 54.5
```

Listing 2: Hierarchical FOMC fit with constant variance

```
saemix version used for fitting:
                                          3.2
mkin version used for pre-fitting: 1.2.4
Date of fit: Fri Aug 4 12:14:23 2023
Date of summary: Fri Aug 4 12:23:55 2023
Equations:
d_{meso}/dt = - (alpha/beta) * 1/((time/beta) + 1) * meso
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 5.212 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
   meso_0 log_alpha log_beta
  93.0520
             0.6008 3.4176
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
         meso_0 log_alpha log_beta
meso_0
                     1.53
            6.287
log_alpha 0.000
log_beta 0.000
                        0.00
                                  1.724
Starting values for error model parameters:
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  787.4 793.6 -386.7
Optimised parameters:
                 est.
                            lower
                                     upper
meso_0
               93.5648 91.42864 95.7009

    0.7645
    0.28068
    1.2484

    3.6597
    3.05999
    4.2594

    5.0708
    4.29823
    5.8435

log_alpha
log_beta
a.1
SD.meso_0 0.1691 -34.01517 34.3535
SD.log_alpha 0.3764 0.05834 0.6945
SD.log_beta 0.3903 -0.06074 0.8414
Correlation:
           meso_0 log_lph
log_alpha -0.2839
log_beta -0.3443 0.8855
Random effects:
                est.
                          lower
SD.meso_0 0.1691 -34.01517 34.3535
SD.log_alpha 0.3764 0.05834 0.6945
SD.log_beta 0.3903 -0.06074 0.8414
Variance model:
est. lower upper
a.1 5.071 4.298 5.843
Backtransformed parameters:
         est. lower upper
meso_0 93.565 91.429 95.701
alpha 2.148 1.324 3.485
beta 38.850 21.327 70.770
Estimated disappearance times:
    DT50 DT90 DT50back
meso 14.8 74.64 22.47
```

Listing 3: Hierarchical DFOP fit with constant variance

```
saemix version used for fitting:
                                       3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting: 4.3.1
Date of fit: Fri Aug 4 12:14:23 2023
Date of summary: Fri Aug 4 12:23:55 2023
* meso
Data:
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 11.27 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
  meso_0 log_k1 log_k2 g_qlogis
93.14689 -2.05241 -3.53079 -0.09522
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
        meso_0 log_k1 log_k2 g_qlogis
meso_0
          6.418 0.000 0.000
                                    0.00
        0.000 1.018 0.000
0.000 0.000 1.694
log_k1
log_k2
                                    0.00
g_qlogis 0.000 0.000 0.000
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  787.6 795.6 -384.8
Optimised parameters:
                         lower
               est.
                                 upper
            93.6684 91.63599 95.7008
meso 0
            -1.7354 -2.61433 -0.8565
-3.4015 -3.73323 -3.0697
log_k1
log_k2
           -1.6341 -2.66133 -0.6069
4.7803 4.01269 5.5479
g_qlogis
            0.1661 -30.97086 31.3031
0.1127 -2.59680 2.8223
SD.meso_0
SD.log_k1
SD.log_k2 0.6394 0.41499 0.8638
SD.g_qlogis 0.8166 0.09785 1.5353
Correlation:
         meso_0 log_k1 log_k2
         0.1757
log_k1
log_k2 0.0199 0.2990
g_qlogis 0.0813 -0.7431 -0.3826
Random effects:
             est.
                        lower upper
SD.meso_0 0.1661 -30.97086 31.3031
SD.log_k1 0.1127 -2.59680 2.8223
SD.log_k2 0.6394 0.41499 0.8638
SD.g_qlogis 0.8166 0.09785 1.5353
Variance model:
    est. lower upper
a.1 4.78 4.013 5.548
Backtransformed parameters:
           est.
                   lower upper
meso_0 93.66841 91.63599 95.70082
k1
        0.17633 0.07322 0.42466
        0.03332 0.02392 0.04643
g
        0.16327 0.06529 0.35277
Estimated disappearance times:
```

DT50 DT90 DT50back DT50_k1 DT50_k2 meso 16.04 63.75 19.19 3.931 20.8

Listing 4: Hierarchical SFORB fit with constant variance

```
3.2
saemix version used for fitting:
mkin version used for pre-fitting: 1.2.4
R version used for fitting: 4.3.1
Date of fit: Fri Aug 4 12:14:23 2023
Date of summary: Fri Aug 4 12:23:55 2023
d_meso_free/dt = - k_meso_free * meso_free - k_meso_free_bound *
              meso_free + k_meso_bound_free * meso_bound
d_meso_bound/dt = + k_meso_free_bound * meso_free - k_meso_bound_free *
              meso bound
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 11.191 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
            meso_free_0 log_k_meso_free_log_k_meso_free_bound
                    93.147
                                                -2.305
log_k_meso_bound_free
                    -3.761
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
                           meso_free_0 log_k_meso_free log_k_meso_free_bound
meso_free_0
                                     6.418
                                                     0.0000
                                     0.000
                                                        0.9276
log_k_meso_free
log_k_meso_free_bound
                                 0.000
                                     0.000
                                                        0.0000
                                                                                       2.272
log_k_meso_bound_free
                                                       0.0000
                                                                                       0.000
                          log_k_meso_bound_free
                                                  0.000
meso_free_0
log_k_meso_free
                                                  0.000
log_k_meso_free_bound
                                                  0.000
log_k_meso_bound_free
                                                  1.447
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
  AIC BIC logLik
787.4 795.4 -384.7
{\tt Optimised\ parameters:}
                                est. lower upper 93.6285 91.6262 95.631
meso_free 0
log_k_meso_free

        log_k_meso_iree_bound
        -3.2213
        -4.4695
        -1.973

        log_k_meso_bound_free
        -2.4246
        -3.5668
        -1.282

        a.1
        4.7372
        3.9542
        5.520

        SD.meso_free_0
        0.1634
        -32.7769
        33.104

        SD.log_k_meso_free
        0.4885
        0.3080
        0.669

        SD.log_k_meso_free_bound
        0.2876
        -1.7955
        2.371

        SD.log_k_meso_bound_free
        0.9942
        0.2181
        1.770

Correlation:
                             ms_fr_0 lg_k_m_ lg_k_ms_f_
log_k_meso_free
                             0.2332
log_k_meso_free_bound 0.1100 0.5964
log_k_meso_bound_free -0.0413  0.3697  0.8025
Random effects:
                                   est.
                                             lower upper
SD.meso_free_0
                                0.1634 -32.7769 33.104
SD.log_k_meso_free_bound 0.2876 -1.7955 2.371
SD.log_k_meso_bound_free 0.9942 0.2181 1.770
Variance model:
      est. lower upper
 a.1 4.737 3.954 5.52
```

Listing 5: Hierarchical HS fit with constant variance

```
saemix version used for fitting: 3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
                                      4.3.1
Date of fit: Fri Aug 4 12:14:23 2023
Date of summary: Fri Aug 4 12:23:55 2023
Equations:
d_{meso}/dt = - ifelse(time <= tb, k1, k2) * meso
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 10.613 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
meso_0 log_k1 log_k2 log_tb
92.920 -2.409 -3.295 2.471
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
      meso_0 log_k1 log_k2 log_tb
meso_0 6.477 0.0000 0.0000
log_k1 0.000 0.8675 0.0000
log_k2 0.000 0.0000 0.4035
                                0.00
log_tb 0.000 0.0000 0.0000
Starting values for error model parameters:
a.1
 1
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  781.9 789.9
Optimised parameters:
                       lower
              est.
                               upper
         93.34242 91.4730 95.2118
meso 0
log_k1
          -2.77312 -3.0826 -2.4637
         -3.61854 -3.8430 -3.3941
log_k2
         2.00266 1.3357 2.6696
4.47693 3.7059 5.2479
log_tb
a.1
SD.meso_0 0.07963 -63.1661 63.3253
SD.log_k1 0.47817 0.2467 0.7097
SD.log_k2 0.39216 0.2137 0.5706
SD.log_tb 0.94683 0.4208 1.4728
Correlation:
\begin{array}{cccc} & \texttt{meso\_0} & \texttt{log\_k1} & \texttt{log\_k2} \\ \texttt{log\_k1} & \texttt{0.1627} \end{array}
log_k2 0.0063 -0.0301
log_tb 0.0083 -0.3931 -0.1225
Random effects:
                      lower upper
             est.
SD.meso_0 0.07963 -63.1661 63.3253
SD.log_k1 0.47817 0.2467 0.7097
SD.log_k2 0.39216 0.2137 0.5706
Variance model:
     est. lower upper
a.1 4.477 3.706 5.248
Backtransformed parameters:
            est.
                   lower upper
meso_0 93.34242 91.47303 95.21181
k1
       0.06247 0.04584 0.08512
k2
        0.02682 0.02143 0.03357
tb
        7.40872 3.80282 14.43376
Estimated disappearance times:
    DT50 DT90 DT50back DT50_k1 DT50_k2
meso 16 76 22.88 11.1 25.84
```

Fits with covariate effects

Listing 6: Hierarchichal SFO fit with pH influence

```
saemix version used for fitting: 3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
Date of fit: Fri Aug 4 12:15:27 2023
Date of summary: Fri Aug 4 12:23:55 2023
Equations:
d_{meso}/dt = - k_{meso} * meso
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 7.431 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
    meso_0 log_k_meso
    90.832
              -3.192
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
         meso_0 log_k_meso
meso O
                      0.0000
            6.752
log_k_meso 0.000
                      0.9155
Starting values for error model parameters:  \\
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  783.1 787.5 -386.5
Optimised parameters:
                       est. lower
meso_0 91.3481 89.2688 93.4275 log_k_meso -6.6614 -7.9715 -5.3514 beta_pH(log_k_meso) 0.5871 0.3684 0.8059
               5.4750 4.7085 6.2415
SD.log_k_meso
                     0.3471 0.2258 0.4684
Correlation:
                   meso_0 lg_k_ms
log_k_meso
                     0.0414
beta_pH(log_k_meso) -0.0183 -0.9917
                est. lower upper
SD.log_k_meso 0.3471 0.2258 0.4684
Variance model:
     est. lower upper
a.1 5.475 4.709 6.242
Backtransformed parameters:
            est.
                     lower
                                upper
meso_0 91.348139 8.927e+01 93.427476
k_meso 0.001279 3.452e-04 0.004741
Covariates used for endpoints below:
pH
50% 5.75
Estimated disappearance times:
     DT50 DT90
meso 18.52 61.52
```

Listing 7: Hierarchichal FOMC fit with pH influence

```
saemix version used for fitting: 3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
Date of fit: Fri Aug 4 12:15:33 2023
Date of summary: Fri Aug 4 12:23:55 2023
Equations:
d_{meso}/dt = - (alpha/beta) * 1/((time/beta) + 1) * meso
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 3.962 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
  meso_0 log_alpha log_beta
  93.0520
            0.6008 3.4176
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
        meso_0 log_alpha log_beta
meso_0
           6.287
                      0.00
                    1.53
log_alpha 0.000
log_beta 0.000
                      0.00
                               1.724
Starting values for error model parameters:
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  770.1 776.3 -378
Optimised parameters:
                        est.
                                  lower
                                          upper
                  92.840646 90.750461 94.9308
meso_0
log_alpha
                   -2.206602 -3.494546 -0.9187
beta_pH(log_alpha) 0.577505 0.369805 0.7852
log_beta 4.214099 3.438851 4.9893
                    5.027768 4.322028 5.7335
a.1
                    0.004034 -23.766993 23.7751
SD.log_alpha
                   0.374640 0.009252 0.7400
SD.log_beta
Correlation:
                  meso_0 log_lph bt_H(_)
log_alpha
                   -0.0865
beta_pH(log_alpha) -0.0789 -0.8704
                   -0.3544 0.3302 0.1628
log_beta
Random effects:
                           lower upper
                 est.
SD.log_alpha 0.004034 -23.766993 23.78
SD.log_beta 0.374640 0.009252 0.74
Variance model:
     est. lower upper
a.1 5.028 4.322 5.734
Backtransformed parameters:
          est.
                 lower
meso_0 92.8406 90.75046 94.9308
alpha 0.1101 0.03036 0.3991
beta 67.6332 31.15113 146.8404
Covariates used for endpoints below:
50% 5.75
Estimated disappearance times:
    DT50 DT90 DT50back
meso 17.28 76.37
```

Listing 8: Refined hierarchical FOMC fit with pH influence

```
saemix version used for fitting:
                                     3.2
mkin version used for pre-fitting: 1.2.4
Date of fur. Fri Aug 4 12:15:44 2023
Date of summary: Fri Aug 4 12:23:55 2023
Equations:
d_{meso}/dt = - (alpha/beta) * 1/((time/beta) + 1) * meso
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 9.31 s
Using 300, 100 iterations and 3 chains \,
Variance model: Constant variance
Starting values for degradation parameters:
  meso_0 log_alpha log_beta
  93.0520
           0.6008 3.4176
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
        meso_0 log_alpha log_beta
meso_0
           6.287
                      0.00
                   1.53
log_alpha 0.000
log_beta 0.000
                     0.00
                              1.724
Starting values for error model parameters:
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  767.5 772.8 -377.7
Optimised parameters:
                     est.
                             lower
                                     upper
                  93.0536 90.9771 95.1300
meso_0
log_alpha
                   -2.9054 -4.1803 -1.6304
beta_pH(log_alpha) 0.6590 0.4437 0.8744
            3.9549 3.2860 4.6239
log_beta
                   4.9784 4.2815 5.6754
a.1
                  0.4019 0.2632 0.5406
SD.log_beta
Correlation:
                  meso_0 log_lph bt_H(_)
                   -0.0397
log_alpha
beta_pH(log_alpha) -0.0899 -0.9146
                   -0.3473 0.2038 0.1919
log_beta
Random effects:
              est. lower upper
SD.log_beta 0.4019 0.2632 0.5406
Variance model:
    est. lower upper
a.1 4.978 4.281 5.675
{\tt Backtransformed\ parameters:}
           est.
                  lower
meso_0 93.05359 90.97713 95.1300
alpha 0.05473 0.01529 0.1958
beta 52.19251 26.73597 101.8874
Covariates used for endpoints below:
50% 5.75
Estimated disappearance times:
    DT50 DT90 DT50back
meso 17.3 82.91
```

Listing 9: Hierarchichal DFOP fit with pH influence

```
saemix version used for fitting: 3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
Date of fit: Fri Aug 4 12:15:52 2023
Date of summary: Fri Aug 4 12:23:55 2023
* meso
Data:
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 7.062 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
  meso_0 log_k1 log_k2 g_qlogis
93.14689 -2.05241 -3.53079 -0.09522
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
        meso_0 log_k1 log_k2 g_qlogis
meso_0
         6.418 0.000 0.000
                                0.00
log_k1
        0.000 1.018 0.000
log_k2
        0.000 0.000 1.694
                                0.00
g_qlogis 0.000 0.000 0.000
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  769.1 777.1 -375.5
Optimised parameters:
                      est.
                              lower
                                      upper
                 92.843344 90.8464 94.84028
meso 0
                  -2.815685 -3.0888 -2.54261
log k1
                -11.479779 -15.3203 -7.63923
log_k2
beta_pH(log_k2) 1.308417 0.6948 1.92203
g_qlogis 3.133036 0.4657 5.80035
g_qlogis
beta_pH(g_qlogis) -0.565988 -1.0394 -0.09262
a.1 4.955518 4.2597 5.65135
                  SD.log_k2
{\tt SD.g\_qlogis}
Correlation:
                meso_0 log_k1 log_k2 b_H(_2) g_qlogs
log_k1
                 0.2706
                 -0.0571 0.1096
log_k2
beta_pH(log_k2) 0.0554 -0.1291 -0.9937
Random effects:
               est.
                    lower upper
SD.log_k2 0.758963 0.4685 1.049
SD.g_qlogis 0.005215 -9.9561 9.967
Variance model:
    est. lower upper
a.1 4.956 4.26 5.651
Backtransformed parameters:
          est.
                  lower
meso_0 9.284e+01 9.085e+01 9.484e+01
k1
     5.986e-02 4.556e-02 7.866e-02
      1.034e-05 2.221e-07 4.812e-04
      9.582e-01 6.144e-01 9.970e-01
g
Covariates used for endpoints below:
```

pH 50% 5.75

Listing 10: Refined hierarchical DFOP fit with pH influence

```
saemix version used for fitting:
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
                                  4.3.1
Date of fit: Fri Aug 4 12:16:05 2023
Date of summary: Fri Aug 4 12:23:55 2023
* meso
Data:
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 10.981 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
  meso_0 log_k1 log_k2 g_qlogis
93.14689 -2.05241 -3.53079 -0.09522
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
        meso_0 log_k1 log_k2 g_qlogis
meso_0
         6.418 0.000 0.000
                                 0.00
        0.000 1.018 0.000
log_k1
log_k2
        0.000 0.000 1.694
                                 0.00
g_qlogis 0.000 0.000 0.000
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  765.1 772.3 -374.6
Optimised parameters:
                    est.
                            lower
                                     upper
                 93.3333 91.2427 95.42394
meso 0
log_k1
                 -1.7997 -2.9124 -0.68698
                 -8.1810 -10.1819 -6.18008
log_k2
beta_pH(log_k2) 0.8064 0.4903 1.12257
g_qlogis 3.3513 -1.1792 7.88182
g_qlogis
beta_pH(g_qlogis) -0.8672 -1.7661 0.03177
a.1 4.9158 4.2277 5.60390
                 0.3946 0.2565 0.53281
SD.log_k2
Correlation:
                 meso_0 log_k1 log_k2 b_H(_2) g_qlogs
                 0.1730
log_k1
log_k2
                  0.0442 0.5370
beta_pH(log_k2) -0.0392 -0.4880 -0.9923
                 -0.1536 0.1431 -0.1129 0.1432
g_qlogis
beta_pH(g_qlogis) 0.1504 -0.3151 -0.0196 -0.0212 -0.9798
Random effects:
           est. lower upper
SD.log_k2 0.3946 0.2565 0.5328
Variance model:
     est. lower upper
a.1 4.916 4.228 5.604
Backtransformed parameters:
           est.
                  lower
meso_0 9.333e+01 9.124e+01 95.42394
k1
      1.654e-01 5.435e-02 0.50309
k2
       2.799e-04 3.785e-05 0.00207
       9.661e-01 2.352e-01 0.99962
g
Covariates used for endpoints below:
     pН
50% 5.75
```

Estimated disappearance times:

DT50 DT90 DT50back DT50_k1 DT50_k2
meso 18.37 73.52 22.13 4.192 23.99

Listing 11: Further refined hierarchical DFOP fit with pH influence

```
saemix version used for fitting:
                                        3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
                                      4.3.1
Date of fit: Fri Aug 4 12:16:20 2023
Date of summary: Fri Aug 4 12:23:55 2023
* meso
Data:
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 7.515 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
  meso_0 log_k1 log_k2 g_qlogis
93.14689 -2.05241 -3.53079 -0.09522
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
         meso_0 log_k1 log_k2 g_qlogis
meso_0
          6.418 0.000 0.000
                                     0.00
         0.000 1.018 0.000
log_k1
log_k2
         0.000 0.000 1.694
                                     0.00
g_qlogis 0.000 0.000 0.000
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  767.4 773.6 -376.7
Optimised parameters:
                             lower
                    est.
                                     upper
                 93.3011 91.1905 95.4118
meso 0
log_k1
                 -2.1487 -2.7607 -1.5367
                 -8.1039 -10.4225 -5.7853
log_k2

    log_kz
    -0.1009

    beta_pH(log_k2)
    0.7821

    0.4126
    1.1517

    g_qlogis
    -1.0373
    -1.9337
    -0.1409

    a.1
    5.0095
    4.3082
    5.7108

    SD.log_k2
    0.4622
    0.3009
    0.6235

Correlation:
                 meso_0 log_k1 log_k2 b_H(_2)
log_k1
                 0.2179
                  0.0337 0.5791
log_k2
beta_pH(log_k2) -0.0326 -0.5546 -0.9932
                 0.0237 -0.8479 -0.6571 0.6123
g_qlogis
Random effects:
             est. lower upper
SD.log_k2 0.4622 0.3009 0.6235
Variance model:
     est. lower upper
a.1 5.009 4.308 5.711
Backtransformed parameters:
            est.
                     lower
meso_0 9.330e+01 9.119e+01 95.411751
k1
       1.166e-01 6.325e-02 0.215084
       3.024e-04 2.975e-05 0.003072
       2.617e-01 1.263e-01 0.464832
g
Covariates used for endpoints below:
50% 5.75
Estimated disappearance times:
```

DT50 DT90 DT50back DT50_k1 DT50_k2
meso 17.09 73.67 22.18 5.943 25.54

Listing 12: Hierarchichal SFORB fit with pH influence

```
saemix version used for fitting:
                                     3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
                                   4.3.1
Date of fit: Fri Aug 4 12:16:29 2023
Date of summary: Fri Aug 4 12:23:55 2023
d_meso_free/dt = - k_meso_free * meso_free - k_meso_free_bound *
           meso_free + k_meso_bound_free * meso_bound
d_meso_bound/dt = + k_meso_free_bound * meso_free - k_meso_bound_free *
           meso bound
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 6.478 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
          meso_free_0 log_k_meso_free log_k_meso_free_bound
               93.147
                                     -2.305
log_k_meso_bound_free
               -3.761
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
                     meso_free_0 log_k_meso_free log_k_meso_free_bound
meso_free_0
                            6.418
                                          0.0000
                            0.000
                                           0.9276
log_k_meso_free
log_k_meso_free_bound
                            0.000
                                           0.0000
                                                                   2.272
log_k_meso_bound_free
                            0.000
                                           0.0000
                                                                  0.000
                    log_k_meso_bound_free
                                      0.000
meso_free_0
log_k_meso_free
                                      0.000
log_k_meso_free_bound
                                      0.000
log_k_meso_bound_free
                                      1.447
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
  AIC BIC logLik
768.8 776.8 -375.4
Optimised parameters:
                                  est.
                                          lower
                                                  upper
                               93.4204 91.3213 95.5195
meso free 0
                              -5.3742 -6.9366 -3.8117
0.4232 0.1769 0.6695
log_k_meso_free
beta_pH(log_k_meso_free)
log_k_meso_free_bound
                              -3.4889 -4.9243 -2.0535
                               -9.9797 -19.2232 -0.7362
log_k_meso_bound_free
beta_pH(log_k_meso_bound_free) 1.2290 -0.2107 2.6687 a.1 4.9031 4.1795 5.6268
                                SD.log_k_meso_free
SD.log_k_meso_bound_free
Correlation:
                              ms_fr_0 lg_k_m_ b_H(___) lg_k_ms_f_ lg_k_ms_b_
log_k_meso_free
                                0.1493
beta_pH(log_k_meso_free)
                               -0.0930 -0.9854
log_k_meso_free_bound
                               0.2439 0.4621 -0.3492
log_k_meso_bound_free
                                0.2188 0.1292 -0.0339 0.7287
beta_pH(log_k_meso_bound_free) -0.2216 -0.0797 -0.0111 -0.6566
                                                                   -0.9934
Random effects:
                           est. lower upper
                        0.3454 0.2252 0.4656
SD.log_k_meso_free
SD.log_k_meso_bound_free 0.1277 -1.9459 2.2012
Variance model:
     est. lower upper
a.1 4.903 4.18 5.627
```

Listing 13: Refined hierarchical SFORB fit with pH influence

```
saemix version used for fitting:
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
                                    4.3.1
Date of fit: Fri Aug 4 12:16:40 2023
Date of summary: Fri Aug 4 12:23:55 2023
d_meso_free/dt = - k_meso_free * meso_free - k_meso_free_bound *
           meso_free + k_meso_bound_free * meso_bound
d_meso_bound/dt = + k_meso_free_bound * meso_free - k_meso_bound_free *
           meso_bound
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 9.511 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
          meso_free_0
                        log_k_meso_free log_k_meso_free_bound
               93.147
                                     -2.305
log_k_meso_bound_free
               -3.761
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
                     meso_free_0 log_k_meso_free log_k_meso_free_bound
meso_free_0
                            6.418
                                          0.0000
                            0.000
                                           0.9276
log_k_meso_free
log_k_meso_free_bound
                            0.000
                                           0.0000
                                                                   2.272
log_k_meso_bound_free
                            0.000
                                           0.0000
                                                                   0.000
                     log_k_meso_bound_free
                                      0.000
meso_free_0
log_k_meso_free
                                      0.000
log_k_meso_free_bound
                                      0.000
log_k_meso_bound_free
                                      1.447
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
  AIC BIC logLik
770.9 777.2 -378.5
{\tt Optimised\ parameters:}
                            est.
                                  lower
meso free 0
                         93.3196 91.1633 95.4760
                         -6.1460 -7.4306 -4.8614
log_k_meso_free
beta_pH(log_k_meso_free) 0.5435 0.3329 0.7542
log_k_meso_free_bound -3.8001 -5.2027 -2.3975
log_k_meso_bound_free
                         -2.9462 -4.2565 -1.6359
                         5.0825 4.3793 5.7856
a.1
                         0.3338 0.2175 0.4502
SD.log_k_meso_free
Correlation:
                         ms_fr_0 lg_k_m_ b_H(__ lg_k_ms_f_
log_k_meso_free
                         0.1086
{\tt beta\_pH(log\_k\_meso\_free)} \ -0.0426 \ -0.9821
log_k_meso_free_bound 0.2513 0.1717 -0.0409
log_k_meso_bound_free
                          0.1297 0.1171 -0.0139 0.9224
Random effects:
                     est. lower upper
SD.log_k_meso_free 0.3338 0.2175 0.4502
Variance model:
     est. lower upper
a.1 5.082 4.379 5.786
Backtransformed parameters:
                                lower
                  93.319649 9.116e+01 95.47601
                0.002142 5.928e-04 0.00774
k_meso_free
```

Listing 14: Hierarchichal HS fit with pH influence

```
saemix version used for fitting: 3.2
mkin version used for pre-fitting: 1.2.4
R version used for fitting:
Date of fit: Fri Aug 4 12:16:47 2023
Date of summary: Fri Aug 4 12:23:55 2023
Equations:
d_{meso}/dt = - ifelse(time <= tb, k1, k2) * meso
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 5.359 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
meso_0 log_k1 log_k2 log_tb
92.920 -2.409 -3.295 2.471
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
      meso_0 log_k1 log_k2 log_tb
meso_0 6.477 0.0000 0.0000
log_k1 0.000 0.8675 0.0000
log_k2 0.000 0.0000 0.4035
                               0.00
log_tb 0.000 0.0000 0.0000
Starting values for error model parameters:
a.1
 1
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  769.8 779.6 -373.9
Optimised parameters:
                    est.
                           lower
                93.32599 91.4658 95.1862
meso 0
                 -5.81463 -7.2710 -4.3583
log_k1
beta_pH(log_k1) 0.47472 0.2334 0.7160
                -6.79633 -8.7605 -4.8322
log k2
beta_pH(log_k2) 0.54151 0.2124 0.8706
                 3.24674 1.2470 5.2465
log_tb
beta_pH(log_tb) -0.09889 -0.4258 0.2280
           4.49487 3.7766 5.2132
0.37191 0.2370 0.5068
a.1
SD.log_k1
                 0.29210 0.0994 0.4848
0.25353 -0.0664 0.5735
SD.log_k2
SD.log_tb
Correlation:
                {\tt meso\_0} \quad {\tt log\_k1} \quad {\tt b\_H(\_1)} \ {\tt log\_k2} \quad {\tt b\_H(\_2)} \ {\tt log\_tb}
log_k1
                 0.0744
beta_pH(log_k1) -0.0452 -0.9915
                0.0066 -0.0363 0.0376
log k2
beta_pH(log_k2) -0.0071 0.0372 -0.0391 -0.9939
log_tb
                -0.0238 -0.1483 0.1362 -0.3836 0.3696
beta_pH(log_tb) 0.0097 0.1359 -0.1265 0.3736 -0.3653 -0.9905
Random effects:
            est.
                   lower upper
SD.log_k1 0.3719 0.2370 0.5068
SD.log_k2 0.2921 0.0994 0.4848
SD.log_tb 0.2535 -0.0664 0.5735
Variance model:
     est. lower upper
a.1 4.495 3.777 5.213
Backtransformed parameters:
            est.
                     lower
meso_0 93.325994 9.147e+01 9.519e+01
k1
       0.002984 6.954e-04 1.280e-02
        0.001118 1.568e-04 7.969e-03
tb
       25.706437 3.480e+00 1.899e+02
```

Covariates used for endpoints below: $$\operatorname{pH}$$ 50% 5.75

Estimated disappearance times:

DT50 DT90 DT50back DT50_k1 DT50_k2

meso 15.65 79.63 23.97 15.16 27.55

Listing 15: Refined hierarchical HS fit with pH influence

```
saemix version used for fitting:
mkin version used for pre-fitting: 1.2.4
R version used for fitting: 4.3.1
Date of fit: Fri Aug 4 12:16:53 2023
Date of summary: Fri Aug 4 12:23:55 2023
Equations:
d_{meso}/dt = - ifelse(time <= tb, k1, k2) * meso
116 observations of 1 variable(s) grouped in 18 datasets
Model predictions using solution type analytical
Fitted in 5.39 s
Using 300, 100 iterations and 3 chains
Variance model: Constant variance
Starting values for degradation parameters:
meso_0 log_k1 log_k2 log_tb
92.920 -2.409 -3.295 2.471
Fixed degradation parameter values:
Starting values for random effects (square root of initial entries in omega):
      meso_0 log_k1 log_k2 log_tb
meso_0 6.477 0.0000 0.0000
log_k1 0.000 0.8675 0.0000
log_k2 0.000 0.0000 0.4035
                                0.00
log_tb 0.000 0.0000 0.0000
Starting values for error model parameters:
a.1
Results:
Likelihood computed by importance sampling
   AIC BIC logLik
  766.5 775.4 -373.2
Optimised parameters:
                            lower
                    est.
                93.3251 91.49823 95.1520
meso 0
                 -5.6796 -7.08789 -4.2714
log_k1
beta_pH(log_k1) 0.4567 0.22400 0.6894
log_k2 -6.6083 -8.33839 -4.8781
beta_pH(log_k2) 0.4982 0.20644 0.7899
            2.7040 2.33033 3.0777
4.4452 3.73537 5.1551
log_tb
a.1
SD.log_k1
                0.3570 0.22104 0.4930
0.2252 0.01864 0.4318
SD.log_k2
SD.log_tb
                0.5488 0.24560 0.8521
Correlation:
                {\tt meso\_0 \ log\_k1 \ b\_H(\_1) \ log\_k2 \ b\_H(\_2)}
                 0.0740
log_k1
beta_pH(log_k1) -0.0453 -0.9912
log_tb
                -0.0658 -0.1135 0.0913 -0.1500 0.1210
Random effects:
            est. lower upper
SD.log_k1 0.3570 0.22104 0.4930
SD.log_k2 0.2252 0.01864 0.4318
SD.log_tb 0.5488 0.24560 0.8521
Variance model:
     est. lower upper
a.1 4.445 3.735 5.155
Backtransformed parameters:
            est.
                    lower
meso_0 93.325134 9.150e+01 95.152036
k1
        0.003415 8.352e-04 0.013962
        0.001349 2.392e-04 0.007611
       14.939247 1.028e+01 21.707445
Covariates used for endpoints below:
```

pH 50% 5.75

Session info

R version 4.3.1 (2023-06-16)

Platform: x86_64-pc-linux-gnu (64-bit)

Running under: Debian GNU/Linux 12 (bookworm)

Matrix products: default

BLAS: /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.11.0 LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.11.0

locale:

- [1] LC CTYPE=de DE.UTF-8 LC NUMERIC=C
- [3] LC_TIME=de_DE.UTF-8 LC_COLLATE=de_DE.UTF-8
 [5] LC_MONETARY=de_DE.UTF-8 LC_MESSAGES=de_DE.UTF-8
- [7] LC_PAPER=de_DE.UTF-8 LC_NAME=C
 [9] LC_ADDRESS=C LC_TELEPHONE=C
 [11] LC_MEASUREMENT=de_DE.UTF-8 LC_IDENTIFICATION=C

time zone: Europe/Berlin
tzcode source: system (glibc)

attached base packages:

- [1] parallel stats graphics grDevices utils datasets methods
- [8] base

other attached packages:

- [1] saemix_3.2 npde_3.3 knitr_1.43 mkin_1.2.4

loaded via a namespace (and not attached):

[1]	gtable_0.3.3	dplyr_1.1.2	compiler_4.3.1	<pre>tidyselect_1.2.0</pre>
[5]	tinytex_0.45	<pre>gridExtra_2.3</pre>	scales_1.2.1	yaml_2.3.7
[9]	fastmap_1.1.1	readxl_1.4.2	lattice_0.20-45	ggplot2_3.4.2
[13]	R6_2.5.1	generics_0.1.3	<pre>lmtest_0.9-40</pre>	tibble_3.2.1
[17]	munsell_0.5.0	pillar_1.9.0	rlang_1.1.1	utf8_1.2.3
[21]	xfun_0.39	cli_3.6.1	magrittr_2.0.3	digest_0.6.31
[25]	grid_4.3.1	mclust_6.0.0	lifecycle_1.0.3	nlme_3.1-162
[29]	vctrs_0.6.2	evaluate_0.21	glue_1.6.2	cellranger_1.1.0
[33]	codetools_0.2-19	zoo_1.8-12	fansi_1.0.4	<pre>colorspace_2.1-0</pre>
[37]	tools_4.3.1	pkgconfig_2.0.3	htmltools_0.5.5	

Hardware info

CPU model: Intel(R) Core(TM) i7-4710MQ CPU @ 2.50GHz

MemTotal: 12165632 kB