

Introduction to BLUPF90 software suite

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BLUPF90 software suite

- Collection of software for computations
 - Focus on applications in Breeding and Genetics
- Fortran 90/95
- No GUI (graphical user interface) programs !!!
- Since 1999 by Ignacy Misztal
- First idea: to solve the MME

$$\begin{bmatrix} \mathbf{X'X} & \mathbf{X'Z} \\ \mathbf{Z'X} & \mathbf{Z'Z} + \mathbf{A}^{-1} \frac{\sigma_e^2}{\sigma_a^2} \end{bmatrix} \begin{bmatrix} \hat{\beta} \\ \hat{u} \end{bmatrix} = \begin{bmatrix} \mathbf{X'y} \\ \mathbf{Z'y} \end{bmatrix}$$

- First software: blupf90
- Second idea: variance components estimation
- Developers?

BLUPF90 software developers



Misztal



Shogo Tsuruta



Andres Legarra



lgnacio Aguilar

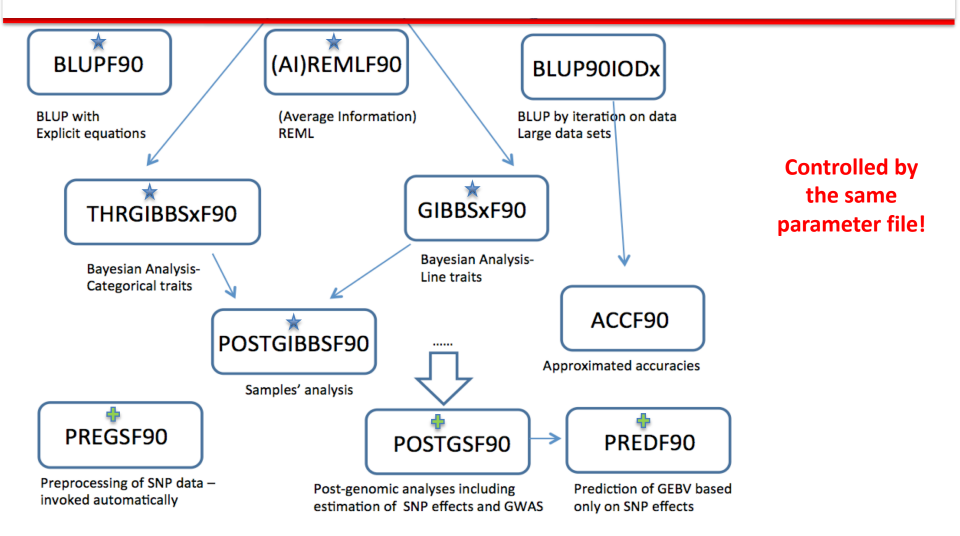


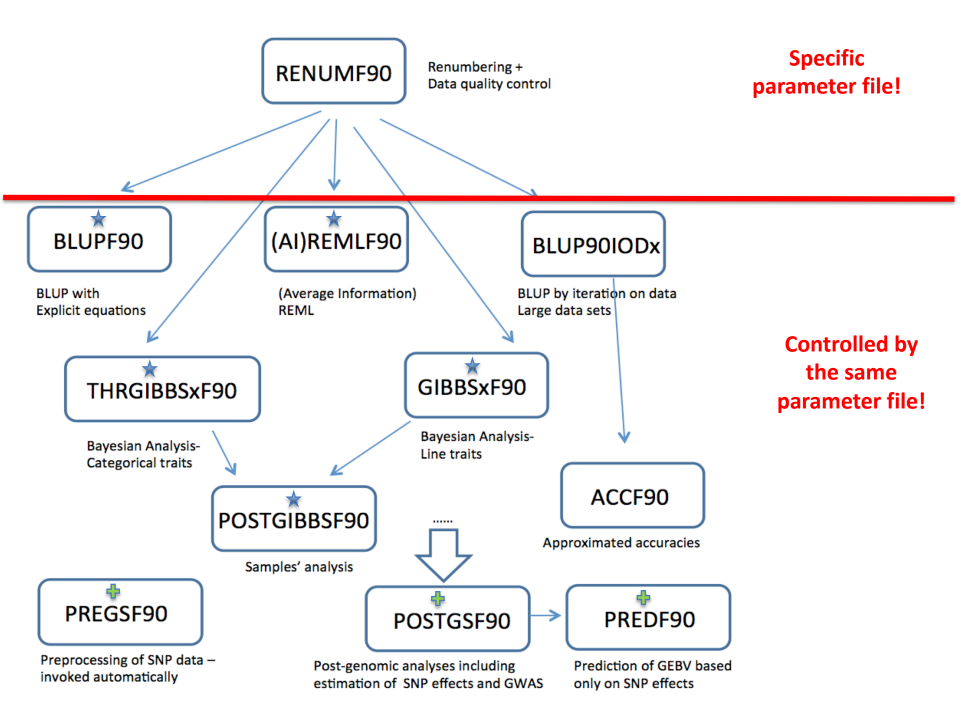
Yutaka Masuda

- + Several contributors
- Research turns into code
- Which programs?

Data File for blupf90 family:

- a) Only numbers Integer or real
- a) All effects need to be renumbered from 1 to N





RENUMF90

A **renumbering program** for the BLUPF90 family of programs

RENUMF90

- Renumbers data and pedigree
- Traces back pedigree for individuals in data file
- Performs comprehensive pedigree checking
- Provides data statistics

RENUMF90

• Supports

- virtually any dataset
- multiple traits
- different effects per trait
- alphanumeric and numeric fields
- unknown parent groups
- covariates for random regression models

RENUMF90 – Input files

- Data file and pedigree file as flat files
 - Columns separated by at least one SPACE
 - No TABS !!!! (current version checks for it)
 - Input files cannot contain character #
 - Missing sire/dams must have code 0
 - code 00 is treated as a known animal

• Parameter file for renumf90

- Has its own parameter file!!!!
- Different from parameter file for other programs!!!!

RENUMF90 – Output files

- Create files to be used by BLUPF90 family
 - renf90.dat data recoded
 - renaddxx.ped renumbered pedigree + statistics
 - renf90.tables cross reference file with renumbered and original effects
 - renf90.par– parameter file

- Based on keywords in capital letter followed by a line(s) with the corresponding data item
- Keywords need to be typed exactly
- Keywords need to be in sequential order!!!
- Lines starting with # are treated as comments and are ignored

RENUMF90 parameter file MANDATORY

Keyword	possible value	description
DATAFILE	character	The name of data file to be processed
TRAITS	integer	Position for observation (trait) in the data file
FIELDS_PASSED TO OUTPUT	integer	Position for the columns in the original data that will be passed to the renumbered data without changes Keep empty if not needed
WEIGHT(S)	integer	The position(s) for weight in the data file Keep empty if not needed
RESIDUAL_VARIANCE	real value(s)	Residual (co)variance
EFFECT	(next slide)	Description of an effect Repeatable

Effects

Keyword	Possible value	effect type	form
EFFECT	CT integer (column where the effect is)	cross	alpha
			numer
enectis		COV	

Keyword (only for covariables)	Possible value	form
NESTED	integer (column where the effect is)	alpha numer

RENUMF90 parameter file Model: y = herd + sex + β age + e

DATAFILE
TRAITS
FIELDS_PASSED TO OUTPUT
WEIGHT(S)
RESIDUAL_VARIANCE
EFFECT

data1.txt

ID	herd	sex	age	phen
ID006	Α	1	1.0	3.0
ID009	Α	2	1.0	2.0
ID012	Α	1	2.0	4.0
ID007	В	2	2.0	6.0
ID010	В	1	1.0	3.0
ID013	В	2	2.0	6.0
ID008	C	1	2.0	6.0
ID011	C	2	1.0	6.0
ID014	C	1	1.0	8.0
ID015	C	2	2.0	4.0

Model: $y = herd + sex + \beta age + e$

DATAFILE data1.txt TRAITS 5 FIELDS_PASS 2 WEIGHT(S)	SED TO OUTPUT
RESIDUAL_V	ARIANCE
	#1 st effect - herd
2 cross alph	а
EFFECT	#2 nd effect - sex
3 cross num	
EFFECT	#3 rd effect - age
4 cov	

Fixed model

data1.txt

ID	herd	sex	age	phen
ID006	Α	1	1.0	3.0
ID009	Α	2	1.0	2.0
ID012	Α	1	2.0	4.0
ID007	В	2	2.0	6.0
ID010	В	1	1.0	3.0
ID013	В	2	2.0	6.0
ID008	C	1	2.0	6.0
ID011	C	2	1.0	6.0
ID014	C	1	1.0	8.0
ID015	C	2	2.0	4.0

Random Effects

Keyword after EFFECT	possible value	description
RANDOM	diagonal	Non-correlated
	animal	Correlation structure among animals
	sire	Not implemented!!!

Keyword	possible value	description
OPTIONAL	ре	Permanent environmental
	mat	Maternal
	mpe	Permanent environmental maternal (only if mat is used)

Random effects file section

Keyword after RANDOM (animal or sire only)	possible value	description
FILE	character	Name of the pedigree file for animal or sire models only

Keyword after FILE (for RANDOM animal only)	possible value	description
FILE_POS	integer	Specifies positions in the pedigree for ani sire dam alternate_dam yob Default: 12300 If maternal effect alternate_dam

Keyword (for RANDOM animal only)	possible value	description
SNP_FILE	character	Optional: If genomic info is to be used Name of the SNP file Format: ID 011122211155152222

Pedigree options

Keyword (for RANDOM animal only)	possible value	description
PED_DEPTH	Integer	Optional Specifies the depth of pedigree search Default = 3 All pedigree = 0
Keyword (for RANDOM animal only)	possible value	description
GEN_INT	Integer min avg max	Optional Specifies min, avg, max generation interval; if yob is present Average used to predict yob of parents
Keyword (for RANDOM animal only)	possible value	description
REC_SEX	Integer	Optional Specifies which parent has records Checks if records are found in specific sex

Unknown Parent Group options

Keyword (for RANDOM animal only)	possible value	description
UPG_TYPE		Optional
	yob 1990 1992	UPG assigned based on yob
	in_pedigrees	Missing parent receives -x x is the UPG number
	internal	UPG assigned by a user-written function (not implemented yet)

Inbreeding option

Keyword (for RANDOM animal only)	possible value	description
INBREEDING		Optional
	pedigree	Calculates inbreeding and saves in the renumbered pedigree file
	file	Reads inbreeding from an external file format: original_ID inbreeding (0 to 1)

Random Regression options

Keyword	possible value	description
RANDOM_REGRESSION		Specifies that random regression should be applied to the random effects
	data	If covariables are in the data
	legendre	Generates legendre polynomials (not implemented)

Keyword	possible value	description
RR_POSITON	Integer	Specifies positions of covariables if RANDOM_REGRESSION type is data

(CO)VARIANCES for Random effects

Keyword	possible value	description
(CO)VARIANCES	real	(co)variance for the animal effect dimension should account for number of traits and random correlated effects

32.79	-7.22	-11.07
-7.22	258.06	87.66
-11.07	87.66	194.34

(CO)VARIANCES structure

• 3 trait (T1-T3) and 2 correlated effects (E1-E2)

			E1			E2	
		T1	T2	Т3	T1	T2	Т3
	T1						
E1	T2						
	Т3						
	T1						
E2	T2						
	Т3						

(CO)VARIANCES for Random effects

Keyword	possible value	description
(CO)VARIANCES	real	(co)variance for non-correlated random effects

Keyword	possible value	description
(CO)VARIANCES_PE	real	(co)variance for the PE effect if present

Keyword	possible value	description
(CO)VARIANCES_MPE	real	(co)variance for the MPE effect if present

Creating interaction between effects

Keyword	possible value	description
COMBINE	integer	Should come before DATAFILE COMBINE 7 2 3 4 Columns 2, 3, 4 are combined into 7 They can be numer or alpha

Keyword	optional	possible values		
COMBINE	optional	definition of new field as a combination of existing fields		
DATAFILE	mandatory	name of raw data file		
TRAITS	mandatory	positions of observations in the raw data file		
FIELDS_PASSED	mandatory	positions of items in the raw data file to be passed to renf90.dat		
WEIGHT(S)	mandatory	positions of weights in the raw data file		
RESIDUAL_VARIANCE	mandatory	residual covariance matrix		
EFFECT	mandatory	effect description		
NESTED	optional	positions of nested covatiates		
RANDOM	optional	declaration of random effect		
OPTIONAL	optional	declaration of MAT, PE, MPE		
FILE	optional	name of raw pedigree file		
FILE_POS	optional	positions of animal ID, sire ID and dam ID		
SNP_FILE	optional	name of SNP marker file		
PED_DEPTH	optional	the maximum generation back from animals with phenotype and/or genotype		
GEN_INT	optional	generation interval to set unknown parent groups (UPG)		
REC_SEX	optional	check if records are found in specific sex		
UPG_TYPE	optional	UPG specification		
INBREEDING	optional	create pedigree file with inbreeding code		
RANDOM_REGRESSION	optional	put covariates for random regressions		
RR_POSITION	optional	positions of covariates for random regressions		
(CO)VARIANCES	optional	covariance components		
(CO)VARIANCES_PE	optional	covariance components for animal PE effects		
(CO)VARIANCES_MPE	optional	covariance components for maternal PE effects		
OPTION	optional	option parameters		

RENUMF90 parameter file Options passed to blupf90

• All lines that begin with keyword OPTION are passed to parameter file renf90.par

This allows automatization of process by using scripts

- For example:
 - OPTION sol se

RENUMF90 parameter file Extra options

The following options can added at the end of the parameter file to redefine parameters used to read the input file:

- the default size of character fields (maximum number of characters in a column - 20)

OPTION alpha_size nn where nn is the new size. - the size of th record length (maximum number of characters in a line - 800) OPTION max_string_readline nn where nn is the new size. - the maximun number of fields (maximum number of columns in a line - 100) OPTION max_field_readline nn

where nn is the number of fields.

- Keyword EFFECT is repeated as many time as effects in the model
- If (CO)VARIANCES for any effect are missing, default matrix with 1.0 in diagonal and 0.1 on offdiagonal will be used
 - WARNING: for EM-REML convergence rate is improved if starting values are too large rather than too small !!!

Model: $y = herd + sex + \beta age + e$

DAIAFILE				
data1.txt				
TRAITS				
5				
FIELDS_PAS	SED TO OUTPUT			
2				
WEIGHT(S)				
. ,				
RESIDUAL	VARIANCE			
1.0 _				
EFFECT	#1 st effect - herd			
2 cross alpha				
RANDOM				
diagonal				
(CO)VARIANCES				
0.5				
EFFECT	#2 nd effect - sex			
3 cross numer				
EFFECT	#3 rd effect - age			
4 cov	5			

What if we want to consider herd as random?

data1.txt

ID	herd	sex	age	phen
ID006	Α	1	1.0	3.0
ID009	Α	2	1.0	2.0
ID012	Α	1	2.0	4.0
ID007	В	2	2.0	6.0
ID010	В	1	1.0	3.0
ID013	В	2	2.0	6.0
ID008	C	1	2.0	6.0
ID011	C	2	1.0	6.0
ID014	C	1	1.0	8.0
ID015	C	2	2.0	4.0

RENUMF90 parameter file <u>Model:</u> y = herd + sex + β age + animal + e

DATAFILE

data1.txt

TRAITS

5

FIELDS_PASSED TO OUTPUT

2

0.5

WEIGHT(S)

RESIDUAL_VARIANCE 1.0

#1st effect - herd EFFECT 2 cross alpha #2nd effect - sex EFFECT 3 cross numer #3rd effect - age EFFECT 4 cov #4th effect - animal EFFECT 1 cross alpha RANDOM animal FILE ped1.txt FILE POS 12300 (CO)VARIANCES

What if we want to consider animal effect as random?

ped1.txt

data1.txt

phen

3.0

2.0

4.0

6.0

3.0

6.0

6.0

6.0

8.0

4.0

ID	Sire	Dam	ID	herd	sex	age
ID006	ID001	ID003	ID006	А	1	1.0
ID009	ID001	ID004	ID009	А	2	1.0
ID012	ID001	ID005	ID012	А	1	2.0
ID007	ID001	ID003	ID007	В	2	2.0
ID010	ID001	ID004	ID010	В	1	1.0
ID013	ID002	ID005	ID013	В	2	2.0
ID008	ID002	ID003	ID008	С	1	2.0
ID011	ID002	ID004	ID011	С	2	1.0
ID014	ID002	ID005	ID014	С	1	1.0
ID015	ID002	ID003	ID015	C	2	2.0

RENUMF90 output files

Pedigree file: renaddxx.ped

Data file: renf90.dat

Parameter file: renf90.par

Renumbering table: renf90.table

RENUMF90 output files Pedigree file: renaddxx.ped

• Structure:

- 1. Animal ID (from 1)
- 2. Parent 1 ID or UPG number for parent 1
- 3. Parent 2 ID or UPG number for parent 2
- 4. 3 minus number of known parents
- 5. Known or estimated year of birth (0 if not provided)
- Number of known parents, if animal has genotype: 10+number of known parents
- 7. Number of records
- 8. Number of progeny as parent 1
- 9. Number of progeny as parent 2
- 10. Original animal ID

RENUMF90 output files Pedigree file: renaddxx.ped

• If option for inbreeding is used: Column 4:

inb/upg code = 4000 / [(1+ms)(1-Fs) + (1+md)(1-Fd)]

ms (md) is 0 if sire (dam) is known, and 1 otherwise Fs (Fd) is the coefficient of inbreeding of sire (dam)

Ex: For an animal with both parents known and F=0 inb/upg code = 2000

RENUMF90 output files renumbering tables: renf90.tables

- For each cross-classified effects

 Original ID, count, consecutive number
- Useful
 - To translate solutions from BLUPF90 program into original alphanumeric values
 - Check counts of records by level

DATAFILE data.test TRATTS 3 4 FIELDS PASSED TO OUTPUT 2 1 # passing alphanumeric WEIGHT(S) RESIDUAL VARIANCE 52 2 4 EFFECT 1 1 cross alpha EFFECT 2 2 cross alpha RANDOM animal OPTIONAL mat mpe pe FILE test.ped (CO)VARIANCES 10 3 2 1 3 11 4 5 2 4 12 6 1 5 6 13.01 (CO)VARIANCES PE 5.3 2.1 2.1 4.85 (CO)VARIANCES_MPE 1.03 .27 .27 .85 EFFECT 5 0 cov NESTED 1 0 alpha EFFECT 6 6 cross alpha RANDOM diagonal

Example of RENUMF90 parameter file

data file - data.test

1 aa 34.5 11 12 zz 3 bb 21.333 22 23 xx 8 cc 23.666 33 34 yy 1 dd 29 44 45 xx 3 aa 30 55 56 yy 5 bb 1234567.890 66 67 zz

pedigree file - test.ped

qq 0 0 aa 0 0 bb qq aa cc qq 0 dd 0 aa

RENUMF90 Printout

```
RENUMF90 version 1.93
name of parameter file? testpar1
datafile:data.test
traits: 3 4
fields passed: 2
                             1
R
  5.000 2.000
 2.000 4.000
Processing effect 1 of type cross
item kind=alpha
Processing effect 2 of type cross
item kind=alpha
Optional maternal effect
Optional maternal permanent environment
Optional permanent environment
pedigree file name "test.ped"
positions of animal, sire, dam, alternate dam and yob 1 2 3 0 0
Reading (CO)VARIANCES: 4 x
                                          4
Reading (CO)VARIANCES_PE: 2 x
                                             2
Reading (CO)VARIANCES MPE: 2 x
                                              2
Processing effect 3 of type cov
item kind=alpha
Processing effect 4 of type cross
item kind=alpha
Maximum size of character fields: 20
Maximum size of record (max_string_readline): 800
Maximum number of fields innput file (max field readline): 100
```

RENUMF90 Printout

table with	5 records	ip ients sorted					
added count Effect group table expanded fr		[:] column 0000 to	-	with records		4	levels
added count Effect group table with added count		[:] column ments sorted	1	with		4	levels
added count Effect group table expanded fr table with added count	rom 10	column 000 to ments sorted	-	with records		4	levels
Effect group table expanded fr wrote statistics	rom 10		_	with records		3	levels
Basic statistics				de is 0)			
Pos Min 3 21.333		Mean 0.20578E+06	SD 0 504005	+06	N 6		
4 11.000				100	6		
5 12.000					6		
Correlation matri	ix						
3 4	5						
3 1.00 0.65							
4 0.65 1.00							
5 0.65 1.00	1.00						
Counts of nonzero							
6	6	6					
6	6 6	6					
0	0	6					

RENUMF90 Printout

```
random effect 2
type:animal
opened output pedigree file "renadd02.ped"
read 5 pedigree records
loaded
               3 parent(s) in round
                                             1
Pedigree checks
Number of animals with records:
                                     4
Number of parents without records:
                                       1
Number of phantom dams:
                      2
Total number of animals:
                          7
random effect 4
type:diag
Wrote parameter file "renf90.par"
Wrote renumbered data "renf90.dat"
```

RENUMF90 output files

data file - renf90.dat

34.5 11 1 3	5 12 1 3 aa 1
21.333 22 2	1 3 23 2 1 bb 3
23.666 33 4	4 7 34 4 2 cc 8
29 44 1 2 3	45 1 1 dd 1
30 55 2 3 5	56 2 2 aa 3
1234567.890	66 3 1 3 67 3 3 bb 5

Pedigree file - renadd02.ped

6	3	1	0	2	2	0	0	bb
0	0	1	0	0	0	2	0	qq
0	3	1	0	1	1	0	0	dd
0	0	1	0	0	0	0	1	D@@0000002
0	0	1	0	0	0	0	1	D@@0000001
0	5	1	0	1	2	0	2	aa
6	7	1	0	2	1	0	0	cc
	0 0 0 0 0	0 0 0 3 0 0 0 0 0 5	0 0 1 0 3 1 0 0 1 0 0 1 0 5 1	0 0 1 0 0 3 1 0 0 0 1 0 0 0 1 0 0 5 1 0	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0 0 1 0 0 0 0 3 1 0 1 1 0 0 1 0 0 0 0 0 1 0 0 0 0 5 1 0 1 2	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

renumbering tables - renf90.tables

```
Effect group 1 of column 1 with 4 levels
Value
        #
             consecutive number
121
322
513
814
Effect group 3 of column 1 with 4 levels
Value
             consecutive number
         #
121
322
513
814
Effect group 4 of column 1 with 3 levels
Value
        # consecutive number
xx 2 1
yy 2 2
zz 2 3
```

parameter file - testpar1

```
# Parameter file for program renf90; it is translated to parameter
# file for BLUPF90 family programs.
DATAFILE
data.test
TRAITS
34
FIELDS PASSED TO OUTPUT
2 1 # passing alphanumeric
WEIGHT(S)
RESIDUAL VARIANCE
5 2
24
EFFECT
1 1 cross alpha
EFFECT
2 2 cross alpha
RANDOM
animal
OPTIONAL
mat mpe pe
FILE
test.ped
(CO)VARIANCES
10 3 2 1
3 11 4 5
2 4 12 6
1 5 6 13.01
(CO)VARIANCES PE
5.3 2.1
2.1 4.85
(CO)VARIANCES MPE
1.03 .27
.27 .85
EFFECT
5 0 cov
NESTED
1 0 alpha
EFFECT
6 6 cross alpha
RANDOM
diagonal
```

new parameter file - renf90.par

```
# BLUPF90 parameter file created by RENF90
DATAFILE
renf90.dat
NUMBER OF TRAITS
          2
NUMBER_OF_EFFECTS
         7
OBSERVATION(S)
  1 2
WEIGHT(S)
EFFECTS: POSITIONS IN DATAFILE NUMBER OF LEVELS TYPE OF EFFECT[EFFECT NESTED]
 3 3
             4 cross
 4 4
             7 cross
 5 5
           7 cross
 5 5
           7 cross
 4 4
             7 cross
 6 0
            4 cov 7 0
 8 8
             3 cross
RANDOM RESIDUAL VALUES
  5.000
             2,000
  2.000
             4,000
 RANDOM GROUP
    2 3
 RANDOM TYPE
 add animal
 FILE
renadd02.ped
(CO)VARIANCES
  10.00
             3.000
                        2.000
                                   1.000
             11.00
                        4.000
                                    5.000
  3.000
                                   6.000
  2.000
             4.000 12.00
             5.000
                        6.000
                                   13.01
  1.000
 RANDOM GROUP
    4
 RANDOM_TYPE
 diagonal
 FILE
(CO)VARIANCES
            0.2700
  1.030
 0.2700
             0.8500
 RANDOM GROUP
    5
 RANDOM_TYPE
 diagonal
 FILE
(CO)VARIANCES
  5.300
             2.100
              4.850
  2.100
 RANDOM GROUP
    7
 RANDOM_TYPE
 diagonal
 FILE
(CO)VARIANCES
  1.000
            0.1000
  0.1000
             1.000
```

renumf90 FAQ

- 1) renumf90 cannot find the data file
- 2) How to include quadratic covariable?
- 3) Error when trying to use covariable
- 4) I want to have original IDs in renf90.dat
- 5) Fixed effects in renf90.dat are different from original

renf90.tables

Check for typos

Column in data file

2 cov numer

FIELDS PASSED TO OUTPUT

renumf90 FAQ

8) When and how to run renumf90?

a) Objective to compare models Run renumf90 ONCE with the most complete model Remove effects from renf90.par

b) Objective to compare non-genomic vs genomic model
 Run renumf90 ONCE with SNP file
 For non-genomic: Remove option for SNP file from renf90.par

c) Objective to mask phenotypes for some animals for validation Run renumf90 ONCE with the complete data Remove animals from renf90.dat

blupf90

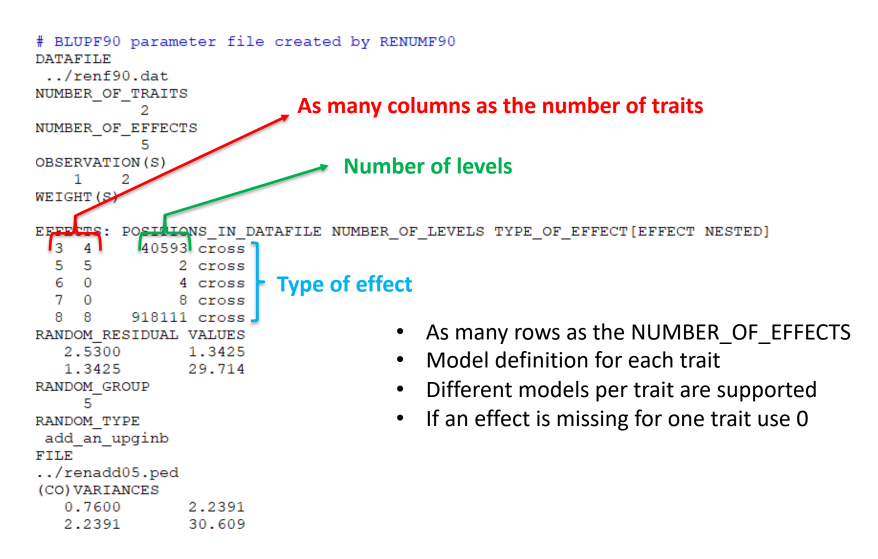
Mixed Model Equations Solver

$$\begin{bmatrix} \mathbf{X'X} & \mathbf{X'Z} \\ \mathbf{Z'X} & \mathbf{Z'Z} + \mathbf{A}^{-1} \frac{\sigma_e^2}{\sigma_a^2} \end{bmatrix} \begin{bmatrix} \hat{\beta} \\ \hat{u} \end{bmatrix} = \begin{bmatrix} \mathbf{X'y} \\ \mathbf{Z'y} \end{bmatrix}$$

blupf90

- Computes generalized solutions by several methods:
 - Preconditioner Conjugate Gradient (PCG)
 - Default Iterative method (fast)
 - Successive over-relaxation (SOR)
 - an iterative method based on Gauss-Seidel
 - Direct solution using sparse Cholesky factorization
 - FSPAK or YAMS (greater memory requirements)
- The solution values change among methods but estimable function should be the same
- Prediction error variances can be obtained using sparse inverse (FSPAK or YAMS)

```
# BLUPF90 parameter file created by RENUMF90
DATAFILE
 ../renf90.dat
NUMBER OF TRAITS
          2
                      Unlimited number of traits and effects
NUMBER OF EFFECTS
          5
OBSERVATION(S)
   1
        2
WEIGHT(S)
EFFECTS: POSITIONS IN DATAFILE NUMBER OF LEVELS TYPE OF EFFECT[EFFECT NESTED]
 3 4
          40593 cross
 5 5
              2 cross
  6 0
             4 cross
 7 0
              8 cross
  8 8 918111 cross
RANDOM RESIDUAL VALUES
  2.5300
          1.3425
  1.3425 29.714
RANDOM GROUP
    5
RANDOM TYPE
add an upginb
FILE
../renadd05.ped
(CO) VARIANCES
  0.7600
               2.2391
         30.609
  2.2391
```



```
# BLUPF90 parameter file created by RENUMF90
DATAFILE
 ../renf90.dat
NUMBER OF TRAITS
NUMBER OF EFFECTS
           5
OBSERVATION(S)
    1
        2
WEIGHT(S)
EFFECTS: POSITIONS IN DATAFILE NUMBER OF LEVELS TYPE OF EFFECT[EFFECT NESTED]
  3 4
           40593 cross
  5 5
               2 cross
  6 0
               4 cross
  7 0
               8 cross
  8
   8 918111 cross
RANDOM RESIDUAL VALUES
                          Should be a square matrix with dimension
                1.3425
   2.5300
                                 equal to the number of traits
   1.3425
                29.714
RANDOM GROUP
     5
RANDOM TYPE

    Use zero (0.0) to indicate uncorrelated residual

add an upginb
FILE
                                   effects between traits
../renadd05.ped

    e.g. For a 3-trait model

(CO) VARIANCES
   0.7600
                2.2391
                                    43.1 0.0 0.0
              30,609
   2.2391
                                    0.0 5.1 3.2
                                    0.0 3.2 10.3
```

```
# BLUPF90 parameter file created by RENUMF90
DATAFILE
 ../renf90.dat
NUMBER OF TRAITS
NUMBER OF EFFECTS
          5
OBSERVATION(S)
   1
        2
WEIGHT(S)
EFFECTS: POSITIONS IN DATAFILE NUMBER OF LEVELS TYPE OF EFFECT[EFFECT NESTED]
  3 4
          40593 cross
 5 5
              2 cross
  6 0
              4 cross
              8 cross
 7 0
  8 8 918111 cross
RANDOM RESIDUAL VALUES
  2.5300
              1.3425
          29.714
  1.3425
                             Definition of random effects
RANDOM GROUP
    5
RANDOM TYPE
add an upginb
                             RANDOM GROUP
FILE
                             RANDOM TYPE
../renadd05.ped
(CO) VARIANCES
                             FILE
  0.7600
               2.2391
  2.2391
               30,609
                             (CO) VARIANCES
```

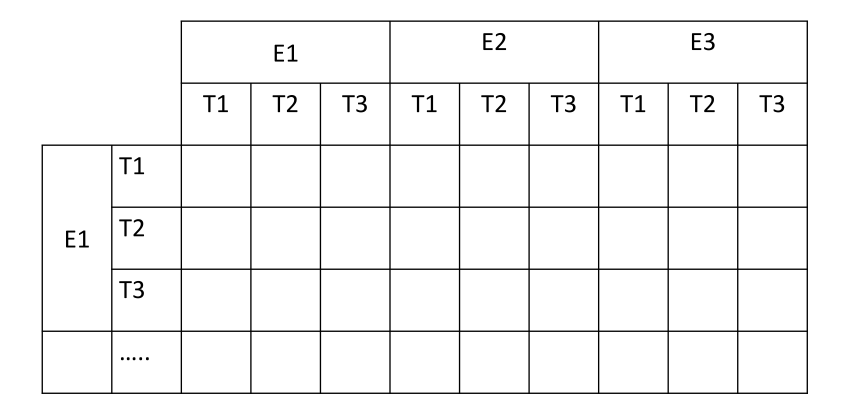
Definition of random effects

• RANDOM_GROUP

- Number(s) of effect from list of effects
- Correlated effects should be consecutive e.g. Maternal effects, Random Regression
- RANDOM_TYPE
 - diagonal, add_animal, add_sire, add_an_upg, add_an_upginb, user_file, user_file_i, or par_domin
- FILE
 - Pedigree file, parental dominance, or user file
- (CO)VARIANCES
 - Square matrix with dimension equal to the number_of_traits*number_of_correlated_effects

(CO)VARIANCES

 Assuming a 3 trait (T1-T3) and 3 correlated effects (E1-E3)



RANDOM_TYPE

- Diagonal
 - for permanent environment effects
 - assumes no correlation between levels of the effect
- add_sire
 - To create a relationship matrix using sire and maternal grandsire
 - Pedigre file:
 - individual number, sire number, maternal grandsire number
- add_animal
 - To create a relationship matrix using sire and dam information
 - Pedigre file:
 - animal number, sire number, dam number

RANDOM_TYPE

- add_an_upg
 - As before but using rules for unknown parent group
 - Pedigre file:
 - animal number, sire number, dam number, parent code
 - missing sire/dam can be replaced by upg number, usually greater than maximum number of animals
 - Parent code = 3 # of known parents
 - 1 both parents known
 - 2 one parent known
 - 3 both parents unknown
- add_an_upginb
 - As before but using rules for unknown parent group and inbreeding
 - Pedigre file:
 - animal number, sire number, dam number, inb/upg code
 - missing sire/dam can be replaced by upg number, usually greater than maximum number of animals
 - inb/upg code = 4000 / [(1+md)(1-Fs) + (1+ms)(1-Fd)]
 - ms (md) is 0 if sire (dam) is known and 1 otherwise
 - Fs(Fs) inbreeding coefficient of the sire (dam)

RANDOM_TYPE

- user_file
 - An inverted matrix is read from file
 - Matrix is stored only upper- or lower-triangular
 - Matrix file:
 - row, col, value
- user_file_i

As before but the matrix will be inverted by the program

• par_domin

A parental dominance file created by program RENDOM

OPTIONS for blupf90

- Programs behavior can be modified by adding extra options at the end of the parameter file
- OPTION option name x1 x2 ...
- option_name: each program has its own definition of options
- The number of optional parameters (x1, x2...) to control the behavior depends on the option

Options for blupf90

Options

OPTION conv_crit 1e-12

Set convergence criteria (deault 1e-12).

OPTION maxrounds 10000

Set maximum number of rounds (default 5000).

OPTION solv_method FSPAK

Selection solutions by FSPAK, SOR or PCG (default PCG).

OPTION r_factor 1.6

Set relaxation factor for SOR (default 1.4).

OPTION sol se

Store solutions and standard errors.

OPTION store_pev_pec 6

Store triangular matrices of standard errors and its covariances for correlated random effects such as direct-maternal effects and randomregression effects in "pev_pec_bf90".

Options for blupf90



OPTION missing -999

Specify missing observations (default 0) in integer.

OPTION residual

y-hat and residual will be included in "yhat_residual".

OPTION blksize 3

Set block size for preconditioner (default 1).

OPTION use_yams

Run the program with YAMS (modified FSPAK).

OPTION SNP_file snp

Specify the SNP file name to use genotype data.

Example of parameter file for blupf90

Single trait "USDA-type" animal model

$$\mathbf{y}_{ijkl} = \mathbf{h}\mathbf{y}\mathbf{s}_i + \mathbf{h}\mathbf{s}_{ij} + \mathbf{p}_k + \mathbf{a}_k + \mathbf{e}_{ijkl}$$

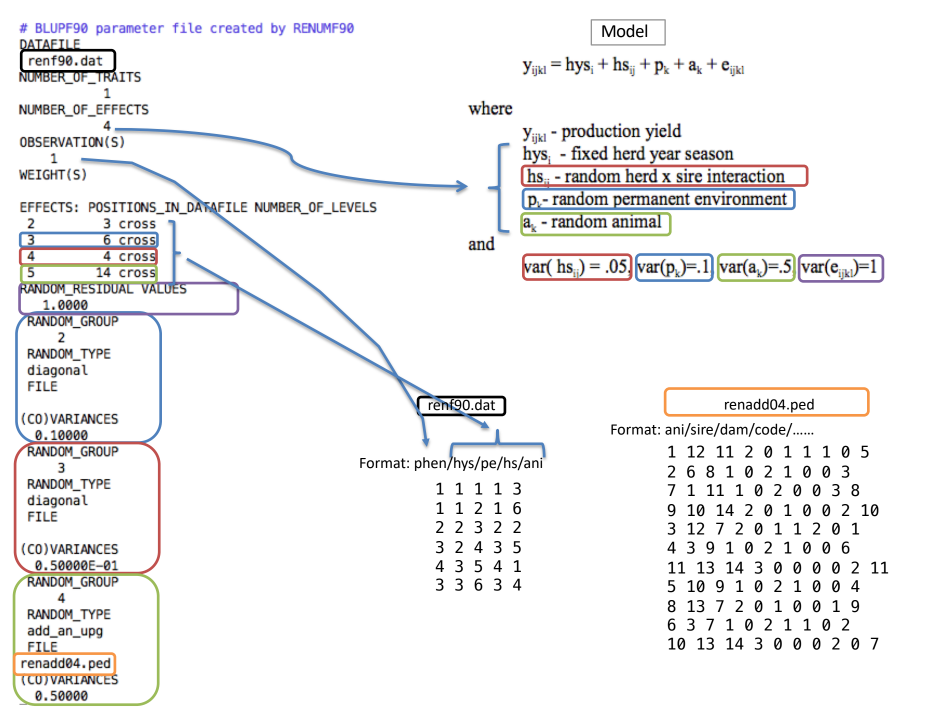
where

 y_{ijkl} - production yield hys_i - fixed herd year season hs_{ij} - random herd x sire interaction p_k - random permanent environment a_k - random animal

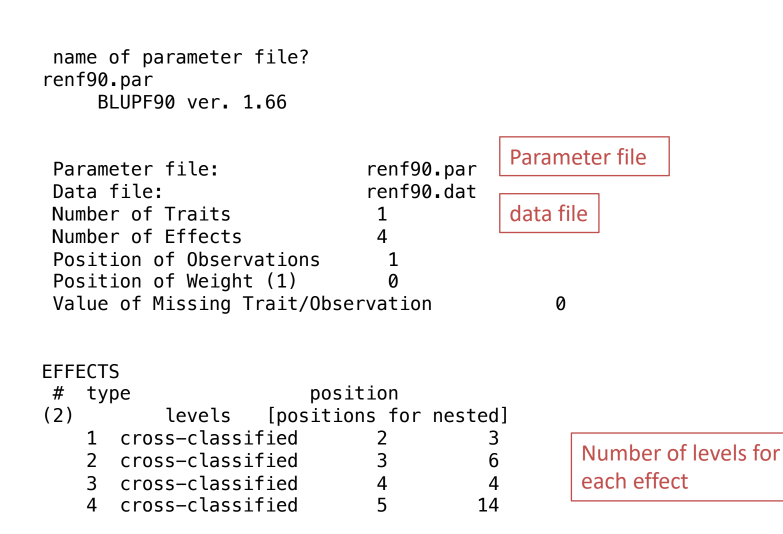
and

$$var(hs_{ij}) = .05, var(p_k) = .1, var(a_k) = .5, var(e_{ijkl}) = 1$$

From blupf90.pdf documentation: http://nce.ads.uga.edu/wiki/doku.php



Output from blupf90



read		6 re	cords in	6.17039	99E-02	s,	Records read from data file
nonzeroes						L	
read				pedigrees	Po	cord	s read from pedigree file
finished	peds	in 6.	1760999E	•	Re	coru	s read from pedigree file
round =	1	converg		0.1435			
round =	2	converg		0.3000E-01			
round =	3	converg		0.1714E-02			
round =	4	converg	ence =	0.2914E-03			
round =	5	converg	ence =	0.1207E-03			
round =	6	converg		0.1278E-03			
round =	7	converg	ence =	0.1085E-03			
round =	8	converg	ence =	0.1405E-03			
round =	9	converg	ence =	0.1904E-03			
round =	10	converg	ence =	0.1119E-03			
round =	11	converg	ence =	0.1562E-04			
round =	12	converg	ence =	0.6141E-05			
round =	13	converg	ence =	0.4609E-05			
round =	14	converg	ence =	0.1750E-04			
round =	15	converg	ence =	0.8708E-04			
round =	16	converg	ence =	0.2800E-03			
round =	17	converg	ence =	0.1286E-04			
round =	18	converg	ence =	0.2309E-06			
round =	19	converg	ence =	0.2566E-08			
round =	20	converg	ence =	0.1131E-09			
round =	21	converg		0.2413E-12			
21 iteı	ratio	ns, co	nvergenc	e criterio	n= 0.24	13E-	12
solutions	s sto	red in f	ile: "so	lutions"	Solutio	ns fil	e

File "solutions"

trait/effect level

solution

	trait	етте/	cτ level	SOLUTION
	(1)	1	1	0.49585171
	1	1	2	2.52240030
$\mathbf{y}_{ijkl} = \mathbf{h}\mathbf{y}\mathbf{s}_i + \mathbf{h}\mathbf{s}_{ij} + \mathbf{p}_k + \mathbf{a}_k + \mathbf{e}_{ijkl}$	$\lfloor 1$	1	3	2.91017217
		Z	1	-0.00965409
where	1	2	2	0.00965406
y _{iikl} - production yield	1	2	3	-0.01930794
hys fixed herd year season	1	2 2	4	0.01930787
hs random herd x sire interaction	1	2	5	0.03861622
$p_{\rm b}$ - random permanent environment	1	2	6	-0.03861599
a _k - random animal		3	1	-0.0000001
	1	3	2	-0.00965384
Parameter File	1	3	3	-0.00965406
DATAFILE	1	3	4	0.01930810
renf90.dat	1	4	1	0.64574095
NUMBER_OF_TRAITS	1	4	2	-0.30035705
	1	4	3	0.61034316
NUMBER_OF_EFFECTS	1	4	4	0.52426082
	1	4	5	0.27486415
OBSERVATION(S)	1	4	6	0.39795337
1	1	4	7	0.18556405
WEIGHT(S)	1	4	8	-0.90212661
	1	4	9	0.63126003
	1	4	10	-0.17807241
EFFECTS: POSITIONS IN DATAFILE	1	4	11	-0.17807202
$2 \qquad 3 \text{ cross}$	1	4	12	1.25233833
3 6 cross	1	4	13	-1.91741245
4 4 cross	1	4	14	1.51299821
5 14 cross				

Common problem in blupf90

- Wrong data file and pedigree name
 - Program may not stop if file name does not exist
 - Check outputs for data file name and number of records and pedigree read

```
round = 4995 convergence =
                                    NaN
round = 4996 convergence =
                                    NaN
round = 4997 convergence =
                                    NaN
round = 4998 convergence =
                                    NaN
round = 4999 convergence =
                                    NaN
round = 5000
              convergence =
                                    NaN
 5001 iterations, convergence criterion=
                                                 NaN
 solutions stored in file: "solutions"
```

renumf90 FAQ

1) Why solutions are not with original ID?

ask Ignacy Misztal

2) How to match solutions with original ID?

Bash scripting

For trait 1 when animal is effect number 2

\$awk '{ if (\$1==1 && \$2==2) print \$3,\$4}' solutions | sort +0 -1 > sol.temp \$awk '{print \$1,\$10}' renadd02.ped | sort +0 -1 > ids.temp \$join -1 +1 -2 +1 ids.temp sol.temp > renumbered_original_EBV

remlf90 and airemlf90

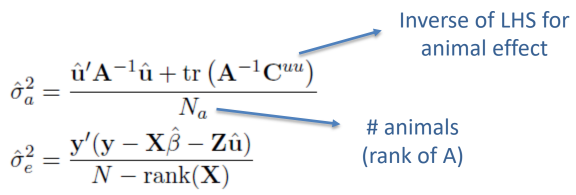
Variance components estimation

$$\begin{bmatrix} \mathbf{X'X} & \mathbf{X'Z} \\ \mathbf{Z'X} & \mathbf{Z'Z} + \mathbf{A}^{-1} \begin{bmatrix} \boldsymbol{\sigma}_{e}^{2} \\ \boldsymbol{\sigma}_{a}^{2} \end{bmatrix} \begin{bmatrix} \hat{\beta} \\ \hat{\boldsymbol{u}} \end{bmatrix} = \begin{bmatrix} \mathbf{X'y} \\ \mathbf{Z'y} \end{bmatrix}$$

remlf90 and airemlf90

- REML = restricted/residual maximum likelihood
 - Patterson and Thompson (1971)
- Most used method to estimate variance components in breeding and genetics
- BLUPF90 family has 2 REML programs
 - remlf90: expectation-maximization (EM) algorithm
 - airemlf90: average information (AI) algorithm
- remlf90 and airemlf90 use the same parameter file as blupf90

remlf90



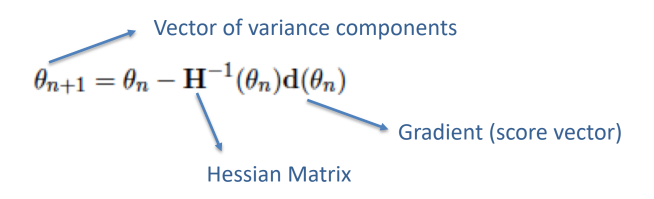
- The equations contain BLUE and BLUP but those values are calculated with known variance components
- This method requires iterations:
- 1. set initial variance components
- 2. compute $\hat{\beta}$ and \hat{u} via mixed model equations
- 3. update and variance components with above equations
- 4. go to 1 or stop if the parameters do not change any more

remlf90

- Simpler equations
- Easier to understand
- More complicated equations in multiple-trait models
- Very slow convergence (looks stable but may not converge)
- Computationally demanding especially for C^{uu}

$$\begin{bmatrix} \hat{\beta} \\ \hat{u} \end{bmatrix} = \begin{bmatrix} \mathbf{X'X} & \mathbf{X'Z} \\ \mathbf{Z'X} & \mathbf{Z'Z} + \mathbf{A}^{-1} \frac{\sigma_e^2}{\sigma_a^2} \end{bmatrix}^{-1} \begin{bmatrix} \mathbf{X'y} \\ \mathbf{Z'y} \end{bmatrix}$$

airemlf90



Average-information algorithm uses this matrix as Hessian,

$$\mathbf{H}(\theta) = \mathcal{I}_A(\theta) = \begin{bmatrix} -\frac{1}{2}\mathbf{y'}\mathbf{PZAZ'PZAZ'Py} & -\frac{1}{2}\mathbf{y'}\mathbf{PZAZ'PPy} \\ -\frac{1}{2}\mathbf{y'}\mathbf{PPZAZ'Py} & -\frac{1}{2}\mathbf{y'}\mathbf{PPPy} \end{bmatrix}$$

Gradient

$$-2\mathbf{d}(\theta) = \begin{bmatrix} \operatorname{tr}(\mathbf{PZAZ'}) - \mathbf{y'PZAZ'Py} \\ \operatorname{tr}(\mathbf{P}) - \mathbf{y'PPy} \end{bmatrix} = \begin{bmatrix} \frac{N_a}{\sigma_a^2} - \frac{\operatorname{tr}(\mathbf{A}^{-1}\mathbf{C}^{uu})}{(\sigma_a^2)^2} - \frac{\hat{\mathbf{u}'A}^{-1}\hat{\mathbf{u}}}{(\sigma_a^2)^2} \\ \frac{N-\operatorname{rank}(\mathbf{X})}{\sigma_e^2} - \frac{1}{\sigma_e^2} \begin{bmatrix} N_a - \frac{\operatorname{tr}(\mathbf{A}^{-1}\mathbf{C}^{uu})}{\sigma_a^2} \end{bmatrix} - \frac{\hat{\mathbf{e}'}\hat{\mathbf{e}}}{(\sigma_e^2)^2} \end{bmatrix}$$

Instructions from Masuda (2018)

Options for remlf90 and airemlf90

OPTION conv_crit 1d-12

Convergence criterion (default 1d-10).

OPTION maxrounds 10000

Maximum rounds (default 5000).

OPTION sol se

Store solutions and se.

OPTION residual

y-hat and residuals will be included in "yhat_residual".

OPTION missing -999

Specify missing observations (default 0) in integer.

OPTION use_yams

Run the program with YAMS (modified FSPAK). The computing time can be dramatically improved.

OPTION constant_var 5 1 2

5: effect number

1: first trait number

2: second trait number

implying the covariance between traits 1 and 2 for effect 5.

Options for airemlf90

OPTION EM-REML 10

Run EM-REML (REMLF90) for first 10 rounds to get initial variances within the parameter space (default 0).

OPTION tol 1d-12

Tolerance (or precision) (default 1d-14) for positive definite matrix and g-inverse subroutines. Convergence may be much faster by changing this value.

OPTION store_pev_pec 6

Store triangular matrices of standard errors and its covariances for correlated random effects such as direct-maternal effects and randomregression effects in "pev_pec_bf90".

Heterogeneous residual variances for a single trait

OPTION hetres_pos 10 11

Specify the column positions of (two) covariables in the data file.

OPTION hetres_pol 4.0 0.1 0.1

Initial values of coefficients for heterogeneous residual variances using In(a0, a1, a2, ...) to make these values.

To transform back to the original scale, use exp(a0+a1*X1+a2*X2)

log-residual function (Foulley and Quaas, 1995)

Options for airemlf90

OPTION se_covar_function <label> <function>

<label>

A name for a particular function (e.g., P1 for phenotypic variance of trait 1, H2_1 for heritability for trait 1, rg12 for genetic correlation between traits 1 and 2, ...).

<function>

A formula to calculate a function of (co)variances to estimate SD. All terms of the function should be written with no spaces.

Each term of the function corresponds to (co)variance elements and could include any random effects (G) and residual (R) (co)variances.

G_eff1_eff2_trt1_trt2

R_trt1_trt1

Examples:

OPTION se_covar_function P G_2_2_1_1+G_2_3_1_1+G_3_3_1_1+G_4_4_1_1+R_1_1

OPTION se_covar_function H2d G_2_2_1_1/(G_2_2_1_1+G_2_3_1_1+G_3_3_1_1+G_4_4_1_1+R_1_1)

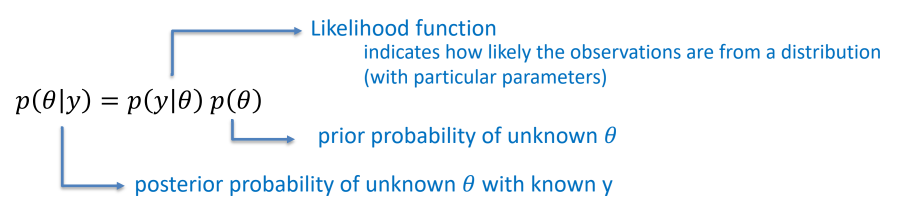
OPTION se_covar_function rg12 G_2_2_1_2/(G_2_2_1_1*G_2_2_2_2)**0.5

Does reml always converge?

- When the expected variance is very small or the covariance matrix is close to non-positive definite, try the following starting values:
 - much smaller = 0.00001
 - much bigger = 1000

- If AIREMLF90 does not converge but REMLF90 converges with the same data set and the same model:
 - run REMLF90 again but with a small starting value to check the estimate because it could be artifact
 - use an option to use EM-REML inside AI-REML: OPTION EM-REML xx where xx is the number of rounds of EM

Bayes Theorem



- Basic idea of Gibbs sampling:
- Gibbs sampling is a numerical method to draw samples from a posterior distribution (not always explicitly available)
- Draw samples = generate random numbers following a distribution
- The results are random numbers (not theoretical formulas)
- The posterior distribution will be drawn based on the numerical values (like a histogram)

Ingredients for Gibbs sampling

1) Theoretical derivation: conditional posterior distribution for each unknown parameter

2) Software: a random number generator for a particular distribution

```
# Basic Gibbs sampling for mu (normal) and sigma2 (inverted chi-square)
y <- C(14,16,18)
N \ll length(y)
n.samples <- 100
mu \ll rep(0, n. samples)
sigma2 <- rep(0,n.samples)
# initial value
mu[1] <- 0
sigma2[1] <- 10
# sampling
for(i in 2:n.samples){
   mu[i] <- rnorm(1, mean=mean(y), sd=sqrt(sigma2[i-1]/N)) # using the most recent sigma2
   df <- N-2
   S <- sum((y-mu[i])^2)
   sigma2[i] <- rinvchisq(1, df=df, scale=S) # using the most recent mu
}
```

Instructions from Masuda (2018)

- gibbs1f90: faster for multiple trait models
- gibbs2f90: better for correlated random effects
- gibbs3f90: for heterogeneous residual variance
- Name of parameter file?
 gibbs1.par
- Number of samples and length of burn-in? samples=10,000 to 100,000; burn-in=0
- Give n to store every n-th sample?

10

gibbs1f90 gibbs1.par --rounds 10000 --burnin 0 --thin 10

- Procedure
 - Run gibbsXf90 to estimate variance components
 - Run postgibbsf90 to process the samples and verify convergence
 - Run gibbsXf90 with new variance components to estimate breeding values (2k to 10k rounds)

OPTION fixed_var all 1 2 3

All solutions and posterior means and SD for effects for effects1, 2, and 3 are stored in "all_solutions" and in "final_solutions" every round using fixed variances. Without numbers, all solutions for all effects are stored.

OPTION fixed_var mean 1 2 3

Posterior means and SD for effects1, 2, and 3 in "final_solutions".

OPTION solution all 1 2 3

Caution: this option will create a huge output solution file when you run many rounds and/or use a large model. All solutions and posterior means and SD for effects1, 2, and 3 are stored in "all_solutions" and in "final_solutions" every round. Without numbers, all solutions for all effects are stored.

OPTION solution mean 1 2 3

Caution: this option will create a huge output solution file when you run many rounds and/or use a large model. Posterior means and SD for effects 1, 2, and 3 in "final_solutions".

OPTION cont 10000

10000 is the number of samples run previously when restarting the program from the last run.

OPTION prior 5 2 -1 5

The (co)variance priors are specified in the parameter file.

Degree of belief for all random effects should be specified using the following structure:

OPTION prior eff1 db1 eff2 db2 ... effn dbn -1 dbres

effx correspond to the effect number and dbx to the degree of belief for this random effect, -1 corresponds to the degree of belief of the residual variance.

In this example 2 is the degree of belief for the 5th effect, and 5 is the degree of belief for the residual.

OPTION seed 123 321

Two seeds for a random number generator can be specified.

OPTION SNP_file snp

Specify the SNP file name to use genotype data.

OPTION se_covar_function <label> <function>

gibbs3f90

OPTION hetres_int col nlev

OPTION hetres_int 5 10

The position "5" to identify the interval in the data file and the number of intervals "10" for heterogeneous residual variances.

gibbs3f90

E

	Parameter file (ex5)
Data (datasire) 1 - HYS 2 - sire 3 - y1 4 - heterogeneous clas 5 - y2 cat datasire	DATAFILE datasire NUMBER_OF_TRAITS NUMBER_OF_EFFECTS OBSERVATION(S) WEIGHT(S) EFFECTS: POSITIONS_IN_DATAFILE 1 1 100 cross 2 2 50 cross RANDOM_RESIDUAL VALUES
6 13 317.55 1 644.26 3 10 280.44 1 563.05 37 1 270.52 5 543.63 53 10 286.43 5 579.84	500 100 100 1000 RANDOM_GROUP RANDOM_TYPE diagonal FILE (CO)VARIANCES 75 10 10 150

OPTION hetres_int 4 5

round 98 209. 416. 416. 828. Residual variance, interval 1 df_r 1997 ee/n 99.4738134864675 101. 202. 202. 412. Residual variance, interval 2 df r 1997 ee/n 146.518188769043 148. 296. 296. 602. Residual variance, interval 3 df r 1997 ee/n 198.183671561078 198. 397. 397. 806. Residual variance, interval 4 df r 1997 ee/n 232.307903786663 228. 455. 455. 917. Residual variance, interval 5 df r 1997 ee/n 301.189371418363 311. 622.

622. 0.126E+04

thrgibbsXf90

- thrgibbs1f90: Gibbs sampler for mixed threshold-linear models Thresholds and variances can be estimated or assumed
- thrgibbs3f90: for heterogeneous residual variance

- Name of parameter file?
 gibbs1.par
- Number of samples and length of burn-in? samples=10,000 to 100,000; burn-in=0
- Give n to store every n-th sample?
 10

thrgibbs1f90 gibbs1.par --rounds 10000 --burnin 0 --thin 10

thrgibbsXf90

Options

OPTION cat 0 0 2 5

"0" indicate that the first and second traits are linear. "2" and "5" indicate that the third and fourth traits are categorical with 2 (binary) and 5 categories.

OPTION save_halfway_samples 5000

The program saves every "5000" samples to restart or recover the job right after the last saved samples. It is useful when the program accidentally stopped.

To restart, add OPTION cont 1 to your parameter file and run thrgibbs1f90 again

OPTION thresholds 0.0 1.0 2.0

Set the fixed thresholds. No need to set 0 for binary traits.

OPTION residual 1

The residual variance can be set to 1 but not necessary for categorical traits more than 2 categories. For binary traits, the residual variance is automatically set to 1, so no need to use this option.

- Basic idea of post Gibbs analysis:
- Summarize and visualize the samples drawn by gibbsXf90
- Confirm if the chain converged
- Find the most probable value = posterior mode as a "point estimate"
- Find the reliability of the estimates = the highest posterior density as a "confidence interval"

- Name of parameter file? gibbs1.par
- Burn-in?

0

- Give n to store every n-th sample? (1 means read all samples)
 10
- input files
 - gibbs_samples, fort.99
- output files
 - "postgibbs_samples"
 - all Gibbs samples for additional post analyses
 - "postmean"
 - posterior means
 - "postsd"
 - posterior standard deviations
 - "postout"

at least > 10 is recommended > 30 may be better

number of independent cycles of Gibbs samples

					*****	Monte	Carlo	E	rror by	Time Series	*****		
Pos.	eff1	eff2	trt1	trt2	MCE	Mean		Н	PD	Effective	Median	Mode	Independent
								Inter	val (95%)	sample size			chain size
1	4	4	1	1	1.362E-02	0.9889	0.	.7788	1.215	70.4	0.9844	0.9861	18
2	4	4	1	2	1.288E-02	1.006		0.777	1.219	84.1	1.006	0.952	18
3	4	4	2	2	1.847E-02	1.66		1.347	1.987	80.3	1.652	1.579	25
4	0	0	1	1	9.530E-03	24.47	:	24.07	24.84	425.6	24.47	24.53	2
5	0	0	1	2	8.253E-03	11.84		11.54	12.18	395.8	11.83	11.82	2
6	0	0	2	2	1.233E-02	30.1	:	29.65	30.58	387.8	30.09	29.97	5

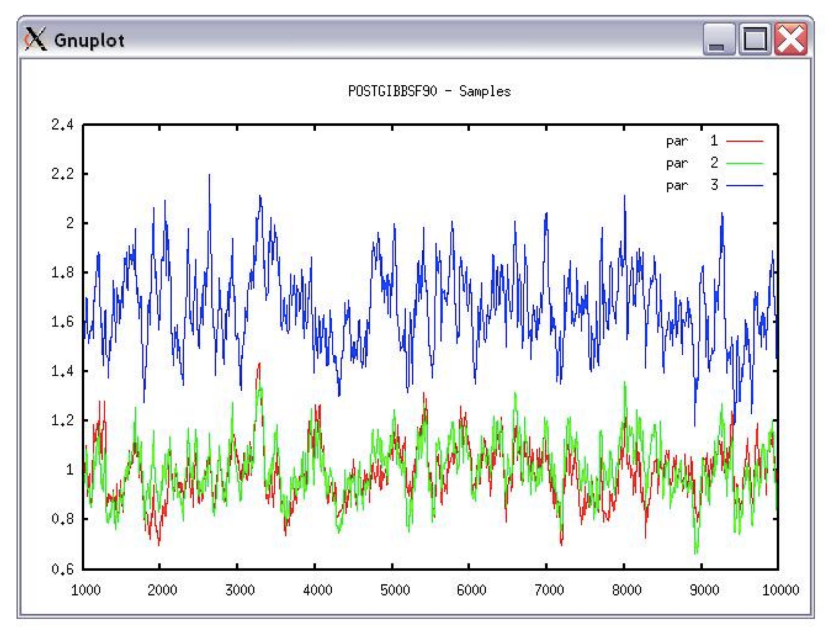
io

ratio between first half and second half of the samples ; should be < 1.0

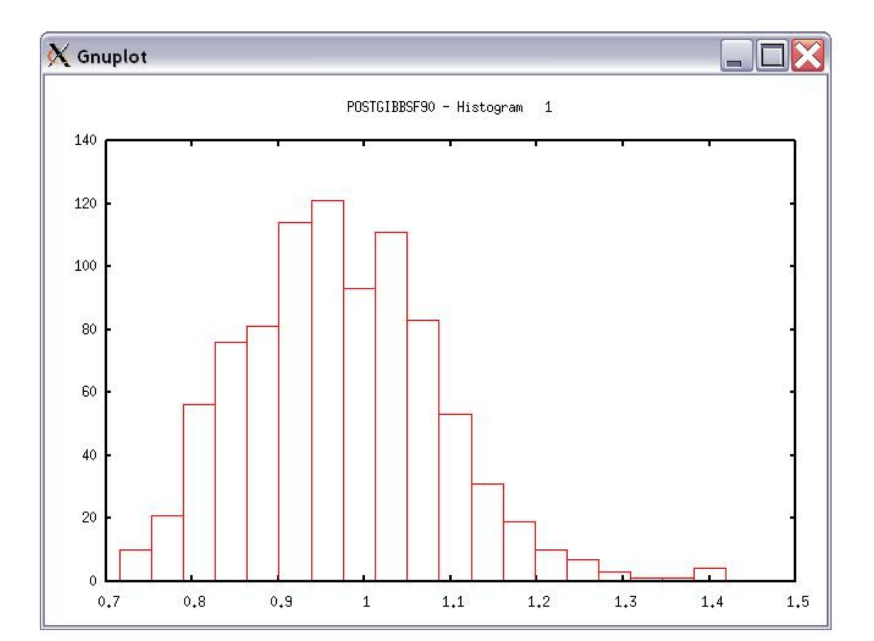
P Lower and upper bounds of Mean ± 1.96PSD

Pos. eff1 eff2 trt1 trt2 PSD		PSD	Mean	PSD		Geweke	Autocorrelations			Independent			
							Interv	7al (95%)	diagnostic	lag: 1	10	50	# batches
1	4	4	1	1	0.1144	0.9889	0.7648	1.213	-0.02	0.853	0.188	0.049	50
2	4	4	1	2	0.1182	1.006	0.7742	1.237	-0.11	0.828	0.111	-0.066	50
3	4	4	2	2	0.1656	1.66	1.335	1.984	0.06	0.828	0.108	-0.021	36
4	0	0	1	1	0.1967	24.47	24.09	24.86	-0.01	0.034	0.029	-0.062	450
5	0	0	1	2	0.1643	11.84	11.51	12.16	0.03	0.032	-0.006	-0.016	450
6	0	0	2	2	0.2429	30.1	29.62	30.57	-0.02	0.07	-0.014	0.037	180

```
Choose a graph for samples (= 1) or histogram (= 2); or exit (= 0)
1
positions
1 2 3 # choose from the position numbers 1 through 6
If the graph is stable (not increasing or decreasing), the convergence is met.
All samples before that point should be discarded as burn-in.
print = 1; other graphs = 2; or stop = 0
2
```



```
Choose a graph for samples (= 1) or histogram (= 2); or exit (= 0)
2
Type position and # bins
1 20
```



Common problems for blupf90 family

- Wrong position or formats for observation and effects
- Misspelling of Keywords
 - Program may stop
- (Co)variance matrices not symmetric, not positive definite
 - Program may not stop
- Large numbers (e.g. 305-day milk yield 10,000 kg)
 - Scale down i.e. 10,000 /1,000 = 10

General output form blupf90 family

- Output printed on the screen is not saved to any file!
- Should use redirection or pipes to store output

renumf90

echo renum.par | renumf90 | tee renum.log

blupf90

echo renf90.par | blupf90 | tee blup.log

airemlf90

echo renf90.par | remlf90 | tee aireml.log

Run in background + Save output

\$bash ai.sh & #can replace bash by sh

```
$vi gibbs.sh
#type the following commands inside ai.sh
    gibbs2f90 <<AA > gibbs.log
    renf90.par
    1000 0
    10
    AA
#save and exit
$bash gibbs.sh & #can replace bash by sh
```