

Pooling estimates of covariance components by penalized maximum likelihood using WOMBAT

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1 Introduction

Combining estimates of covariance components from various sources or multiple analyses of subsets of traits is a common task in quantitative genetics. Specific requirements associated with this task are that the resulting overall matrices should

- be positive (semi-) definite, i.e. not have negative eigenvalues,
- have elements that are ‘similar’ to those from individual analyses,
- not change estimates of variance ratios substantially, and
- result in estimates of phenotypic covariances components that are little distorted.

Single matrices. Most methods in use combine estimates of covariances for a single source of variation at a time; e.g. Mäntysaari (1999) and Thompson et al. (2005).

Recent work has shown that this can result in substantially increased loss in estimates of the corresponding phenotypic covariance matrices, i.e. estimates that are on average further from the population values than unmodified values (Meyer, 2013). The underlying rationale is that pooling individual matrices does not account for sampling correlations between estimates for different sources of variation (from the same part analysis) – and thus does not allow for repartitioning which would keep estimates of the total variation are more or less unchanged.

Multiple matrices. ‘Better’ pooled matrices can be obtained by considering all matrices simultaneously. An early suggestion in this category, termed ‘bending’ (Hayes and Hill, 1981), has been to regress the canonical eigenvalues of the genetic and phenotypic covariance matrices towards their mean (this is often misquoted: regressing the eigenvalues of a single matrix towards their mean is not ‘bending’).

A more flexible alternative is a likelihood based approach which implicitly treats estimates from individual analyses as if they were matrices of corrected sums of squares and cross-products due to some pseudo observations. WOMBAT offers facilities to carry out such calculations as a post-analysis step.

2 Background: The likelihood approach

Consider a vector of observations \mathbf{y} for q traits from a multivariate normal distribution, $\mathbf{y} \sim N(\mathbf{X}\mathbf{b}, \mathbf{V})$, with \mathbf{V} the covariance matrix of \mathbf{y} , \mathbf{b} a vector of fixed effects and \mathbf{X} the corresponding design matrix. If \mathbf{y} can be split into independent parts, \mathbf{y}_i , for instance observations for independent families, \mathbf{V} is block-diagonal. The corresponding log likelihood ($\log \mathcal{L}$) can then be calculated by summing over groups (Thompson, 1976)

$$-2 \log \mathcal{L} \propto \sum_i d_i (\log |\mathbf{V}_i| + \text{tr}(\mathbf{V}_i^{-1} \mathbf{M}_i)) \quad (1)$$

with \mathbf{V}_i the i -th diagonal block of \mathbf{V} , $\mathbf{M}_i = (\mathbf{y}_i - \mathbf{X}_i \hat{\mathbf{b}})(\mathbf{y}_i - \mathbf{X}_i \hat{\mathbf{b}})' / d_i$ the corresponding matrix of corrected mean crossproducts, and d_i denoting the degrees of freedom.

Let Σ_x denote the overall covariance for random effect x for all q traits. For example, for a simple animal model, $x = G, E$ with G and E the genetic and residual covariance matrices, respectively. Further, let \mathbf{S}_i^x be the estimated covariance for random effect x from the i -th part analysis. \mathbf{S}_i^x has expectation Σ_x^x , the submatrix of Σ_x , comprised of the covariance components among the traits represented in the i -th subset of traits. Considering a single source of variation only, estimates of Σ_x could then be obtained by replacing \mathbf{M}_i with \mathbf{S}_i^x and \mathbf{V}_i with Σ_x^x in (Eq. 1) above, and maximising the resulting log likelihood whilst constraining $\hat{\Sigma}_x$ to be positive definite.

2.1 A pseudo pedigree structure

For multiple sources of variation, strong sampling correlations between estimates can be approximated by constructing corresponding terms assuming a simple, balanced pseudo pedigree structure. Let this pseudo structure involves families with m members each, and define matrices \mathbf{C}^x (of size $m \times m$) which give the coefficients for the x -th covariance component in the expectation of covariances between family members. This gives

$$-2 \log \mathcal{L} \propto \sum_i d_i \left(\log \left| \sum_x \mathbf{C}^x \otimes \Sigma_i^x \right| + \text{tr} \left(\left[\sum_x \mathbf{C}^x \otimes \Sigma_i^x \right]^{-1} \left[\sum_x \mathbf{C}^x \otimes \mathbf{S}_i^x \right] \right) \right) \quad (2)$$

with \otimes denoting the direct matrix product. Maximizing (Eq. 2), subject to appropriate constraints, then yields pooled estimates of covariance matrices for all sources of variation.

Suitable choices for the pseudo pedigree depend on the covariance matrices to be pooled – the structure chosen should comprise a sufficient number of types of covariances between relatives so that, when equating these to their expectations, all components can be separated. For a simple animal model, a paternal half-sib design, with m paternal half-sibs per family, suffices. Coefficient matrices for this case are $\mathbf{C}^E = \mathbf{I}_m$ and $\mathbf{C}^G = \frac{1}{4}\mathbf{J}_m + \frac{3}{4}\mathbf{I}_m$ (with \mathbf{J}_m a matrix of size $m \times m$ with all elements equal to unity). If there are common environmental covariances between full-sibs in addition ($x = G, C, E$ with Σ_C the common environmental covariance matrix), a hierarchical full-sib design with n offspring per dam and $k = m/n$ dams per sire would be represented by coefficient matrices $\mathbf{C}^E = \mathbf{I}_m$, $\mathbf{C}^C = \mathbf{I}_k \otimes \mathbf{J}_n$ and $\mathbf{C}^G = \frac{1}{4}\mathbf{J}_m + \mathbf{I}_k \otimes \left(\frac{1}{4}\mathbf{J}_n + \frac{1}{2}\mathbf{I}_n \right)$. For analyses with maternal genetic effects, the pseudo family should include data on at least two generations. e.g. a sire mated to two unrelated dams with two offspring per dam with records on all individuals would provides sire- and dam-offspring as well as full- and half-sib covariances.

2.2 Penalizing the likelihood

In addition, the likelihood approach provides the opportunity for regularized estimation, analogous to that used in standard, multivariate analyses. This involves placing a penalty on the likelihood aimed at reducing sampling variation and thus to improve estimates. In particular, penalties designed to ‘borrow strength’ from estimates of the phenotypic covariance matrix by shrinking estimates of covariance matrices for individual random effects towards their sum or by reducing the spread of estimated canonical eigenvalues, have been shown to yield substantial reductions in sampling variation (Meyer and Kirkpatrick, 2010; Meyer, 2011, 2016).

Corresponding penalties can be employed when pooling estimates from part analyses by maximizing

$$\log \mathcal{L}_P = \log \mathcal{L} - \frac{1}{2} \psi \mathcal{P} \quad (3)$$

instead of $\log \mathcal{L}$, with \mathcal{P} denoting the penalty ($\mathcal{P} > 0$) and ψ a so-called tuning factor ($\psi \geq 0$), determining the emphasis to be given to \mathcal{P} .

3 Pooling using WOMBAT

A facility for pooling estimates of covariance components using the approach described has been implemented in WOMBAT. The main features are:

- Pooling is invoked using the *run time* option `--pool`.
- The *parameter file* should contain a block of statements, framed by `POOL` and `END`, which specify
 - the pseudo-pedigree structure to be used,
 - the type of penalty to be applied and the tuning factor(s) to be used (if any),
 - the minimum eigenvalues of the pooled covariance matrix,
 - the convergence criterion,
 - the form of input files, and
 - whether a ‘minimum’ or ‘full’ parameter file is supplied.
- Input consists of estimates of covariance components from analyses of partial, overlapping subsets of traits:
 - either individual files as generated by WOMBAT or a single, user generated file are accepted,
 - different part analyses can be assigned different weights to be given.
- The implementation allows for multiple random effects, assuming all pooled matrices have the same dimension. Estimates of residual covariances components with a value (or average) of zero are treated as covariances between traits measured on distinct subsets of animals and fixed at that value.

3.1 Pseudo pedigree structures

WOMBAT has several ‘built-in’ pedigree structures and, in addition, allows for other chosen (‘user-defined’) structures by supplying the matrices of coefficients in the expectations of covariances between relatives (matrices \mathbf{C}^x above).

NB: As emphasized above, the structure to be used should be chosen so that there are sufficient types of covariances between relatives to solve for all causal covariances by equating the former to their expectations.

The pseudo pedigree structure is given by a line (within the POOL block) starting with the directive PSEUPED, followed by one of the following options (space separated) and, depending on the option, additional numbers.

PHS selects a balanced, paternal half-sib design comprised of s sires with n progeny each, given by respective integer numbers. If not given, default values of 10 and 4 are used. This design is suitable for a simple animal model.

If the MINPAR option is used, this must be followed by a line with the DIRADD directive.

Example

```

1  VAR  animal  4  NOSTART
2  VAR  residual 4  NOSTART
3  POOL
4      MINPAR
5      PSEUPED PHS 100 5
6      DIRADD  animal
7  END

```

HFS selects a balanced hierarchical full-sib design comprised of s sires, d dams per sire and n progeny per dam. Assumed values for s , d and n can be given (space-separated) on the same line. If omitted, default values of $s = 10$, $d = 5$ and $n = 4$ are used. This design is suitable for a simple animal model or a model fitting maternal permanent environmental effects in addition.

If the MINPAR option is used, codes DIRADD and MATPE need to be given in addition as described above. If not, WOMBAT uses the covariance options given in the MODEL block to identify random effects.

BON selects a design comprising f families of size $n = 8$ due to Bondari et al. (1978). Again, this can be followed by an optional number of families f (default $f = 2$). For this design, expectations of covariances between relatives due to direct and maternal effects are available.

Again, for WOMBAT to 'know' which random effect has which structure, additional information is required. This should comprise one additional line per random effect fitted, with each line consisting of a keyword specifying the type of random effect followed (space separated) by the name of the effect as specified in the VAR statements.

Keywords recognized are:

DIRADD for direct additive genetic,

MATADD for maternal genetic, and

MATPE for maternal permanent environmental effects.

Example

```

1  VAR  animal  6  NOS
2  VAR  gmdam   6  NOS
3  VAR  pedam   6  NOS
4  VAR  residual 6  NOS
5  POOL
6      PSEUPED  BON 10
7      DIRADD  animal
8      MATADD  gmdam
9      MATPE   pedam
10 END

```

In-built variations on Bondari et al. (1978, 's) design can be accessed by specifying options F7, F11 of F17 instead; these should only be required for complicated

models.

USR followed (space separated) by an integer value n and, optionally, by a value f selects a user-defined structure consisting of f families of size n . If not given, a value of $f = 2$ is used. This should be followed by the *upper* triangle of the $n \times n$ matrix of coefficients (\mathbf{C}^x) for all random effects fitted (i.e. excluding residual covariances). For multiple random effects, these should be in the *same* order as the corresponding VAR statements in the parameter file, and each matrix \mathbf{C} should start on a new line.

Example

```

1  VAR  animal  4  NOSTART
2  VAR  residual 4  NOSTART
3  POOL
4      PSEUPED  USR  5
5      1.0  0.50  0.50  0.50  0.50
6      1.0  0.25  0.25  0.25
7      1.0  0.25  0.25
8      1.0  0.25
9      1.0
10  END
```

This shows the coefficients for direct additive genetic effects ('animal') for a family comprising a sire with four progeny from unrelated dams.

3.2 Penalties

WOMBAT first pools estimates of covariance matrices without a penalty on the likelihood. Additional analyses, subject to a penalty are invoked by a line starting with PENALTY followed (space separated) by a code word(s) defining the type and strength of penalty to be applied. Codes for the for the penalty types described in Meyer (2011) recognised are:

CANEIG selects a penalty on the canonical eigenvalues. This has to be followed (space separated) by either ORG or LOG specifying a penalty on eigenvalues on the original scale or transformed to logarithmic scale (no default).

COVARM specifies shrinkage of each covariance matrix towards a given target.

CORREL chooses shrinkage of each correlation matrix towards a given target correlation matrix.

Either COVARM or CORREL can be followed (space separated) by the keyword MAKETAR. If this is given, WOMBAT determines the shrinkage target as the phenotypic covariance (or correlation) matrix obtained by summing estimates of covariances for all sources of variation from the preceding, unpenalized analysis. If this is not given, the upper triangle of the $q \times q$ target matrix is expected to be read from a file with the standard name PenTargetMatrix.

In addition, 'simple' penalties derived assuming a Beta distribution (Meyer, 2016) are available:

KANEIG selects a penalty on canonical eigenvalues.

PACPAC chooses shrinkage of partial (auto-)correlations, either towards zero or towards phenotypic values.

The last entry on the line relates to the tuning factor(s) to be used:

- If a single penalized analysis is to be carried out, the corresponding tuning factor should be given (real value).
- To specify multiple penalized analyses, specify the number of separate tuning

factors multiplied by -1 as an integer value (e.g. -3 for three analyses), and list the corresponding tuning factors space separated on the next line.

Example

1. Shrink all correlation matrices towards the phenotypic correlation matrix using a single tuning factor of 0.1; calculate the shrinkage target from unpenalized results.

```
1  PENALTY CORREL  MAKETAR  0.1
```

2. Shrink canonical eigenvalues on the logarithm scale towards their mean, using 5 different tuning factors

```
1  PENALTY CANEIG LOG  -3
2    0.1  0.5  1.0
```

3. Penalize canonical eigenvalues assuming a Beta distribution with 'sample size of the prior' of 4

```
1  PENALTY KANEIG 4
```

3.3 Other options

WOMBAT recognizes the following other options within the POOL block:

MINPAR The default assumption for the parameter file used is that it is a 'full' parameter file as for a corresponding, multivariate analysis. However, the information on pedigree and data file and their layout is not used. Hence the option MINPAR (on a line by itself) is provided which allows use of a 'minimum' parameter file, switching off some of the consistency checks otherwise carried out. If used, MINPAR must be the *first* entry within the POOL block.

The minimum information to be given in the parameter file must comprise:

1. The ANALysis statement
2. A VAR line for each covariance matrix, together with the NOSTART option telling WOMBAT not to expect a matrix of starting values.
3. The POOL block, including statements showing which random effect represents which type of genetic or non-genetic effect.

Example

```
1  ANAL MUV 14
2  VAR  animal  14  NOSTART
3  VAR  residual 14  NOSTART
4  POOL
5    MINPAR
6    SMALL 0.001d0
7    PSEUPED hfs 100 10 4
8    DIRADD animal
9  END
```

SINGLE The default form of input for results from part analysis is to read estimates from separate files, in the form of output generated by WOMBAT when carrying out multiple analyses of subsets of traits. Alternatively, all information can be given in a single file. This is selected by the option SINGLE, followed (space separated) by the name of the input file (same line).

SMALL followed (space separated) by a real number which gives the lower limit (≤ 1) for smallest eigenvalue allowed in the combined matrices. If this is not specified, a

default value of 0.0001 is used.

DELTA followed (space separated) by a real number specifying the convergence criterion used in the iterative pooling scheme: if the increase in log likelihood between iterates falls below this value, convergence is assumed to have been achieved. If not specified a stringent default of 0.00005 is used.

3.4 Input: Results from partial analyses

3.4.1 Carrying out part analyses using WOMBAT

WOMBAT is set up to allow individual part analyses to be set up and carried out easily:

1. Once the parameter file for the 'full' model (comprising all traits) has been carried out, the parameter files needed for individual analyses can be generated by running WOMBAT with run option `--subset`; see manual for details. In brief, `--subset` will write out parameter files for bivariate analyses for all pairs of traits and provide a (Linux) shell script to carry out the necessary runs in addition. `--subset1` generates corresponding files for univariate analyses and `--subsetn` will allow for selected, n -variate analyses.
2. WOMBAT writes out estimates of covariance components for individual subset analyses to separate files which can directly be used as input for a pooling run.

3.4.2 Option: Input from individual files generated by WOMBAT

The first input option integrates with the facility in WOMBAT to carry out analyses considering selected traits in a multivariate analysis only.

This is invoked by the notation " k " $\rightarrow m$ when specifying the trait numbers in the MODEL block, which stipulates that trait number k in the data file (ranging from 1 to q) should be replaced with value m for the analysis. This notation (generated by `--subset`) causes any records with trait numbers not selected to be skipped and WOMBAT

- writes out the estimates of covariance components together with information on the trait (re)numbering to a file fit the standard name `EstimSubsetk+...+l.dat` with k to l the original trait numbers (in the data file, ranging from 1 to q) for the subset of traits considered in the current analysis.
- appends the name `EstimSubsetk+...+l.dat` to a file named `SubSetsList` followed by a value of 1.00. The latter is a placeholder for the weight to be given to the results in this file, which can be replaced as required.

When encountering the `--pool` option *without* the SINGLE qualifier in the POOL block, WOMBAT tries to acquire the names of the input files and weights for individual analyses from `SubSetsList`. The individual input files listed then must provide the following information:

- (a) The number of traits in the subset (first line).
- (b) The corresponding (original) trait numbers in the 'full' analysis (second line).

This is followed by the covariance matrices estimated. The first matrix given must be the matrix of residual covariances, the other covariance matrices should be given in the same order as specified in the parameter file.

- (c) The first line for each covariance matrix should list the running number of the random effect (0 for 'residual'), the order of fit and the name of the effect (corresponding to a

name in the VAR statement).

- (d) The following lines should give the elements of *full-stored* covariance matrix, with each row of the matrix starting on a new line.

3.4.3 Option: Input from a single, user-generated file

The second option assumes all partial estimates have been collected in a single file. This is specified by the SINGLE option described above.

For each part analysis, this file should contain the following information:

1. A line giving (space separated):
 - a) The number of traits, q_i , in the part analysis ($1 \leq q_i \leq q$, with q the total number of traits).
 - b) The (running) numbers of these traits in the full covariance matrix.
 - c) The relative weight to be given to this part; this can be omitted and, if not given, is set to 1.
2. The elements of the upper triangle of the *residual* covariance matrix, given row-wise, i.e. $q_i(q_i + 1)/2$ elements).
3. For each random effect fitted, the elements of the upper triangle, given row-wise ($q_i(q_i + 1)/2$ elements). Each matrix must begin on a new line and the matrices must given in the same order as the corresponding VAR statements in the parameter file.

3.5 Output

Output from a run with option `--pool` consists of the following files:

1. `PoolEstimates.out` is the main output which summarizes characteristics of the part estimates provided, options chosen, and results for all analyses carried out.
2. `PoolBestPoint` is the equivalent to `BestPoint`, suitable for further examination or as starting values for full, multivariate analyses. The first line gives the likelihood, the number of parameters and tuning parameter used. Then the elements of the upper triangle of the pooled covariance matrices are given, starting with the residual covariance matrix and the other covariance matrices in order of the VAR statements in the parameter file. Each matrix begins on a new line.
If penalized analyses are carried out, copies labelled `PoolBestPoint_unpen` and `PoolBestPoint_txx`, with xx equal to the tuning factor, are generated so that files for all sub-analyses are available at the end of the run.

4 Examples

4.1 Details for simple case with four traits

Example 4 (in the suite of WOMBAT worked examples) gives data for four traits measured by ultra-sound scanning of beef heifers. The model of analysis is a simple animal model. Sub-directory E illustrates pooling of estimates from 6 bivariate analyses.

Results are summarized in the single file:

PartAll.dat		
2 1 2		
3.53021846755401	1.95048936242497	2.08907657452187
2.36692690581371	1.34229728294400	0.977425661859041
2 1 3		
3.55743271408901	2.76836382462327	23.6200689402088

2.32842498547499	0.993394132647659	7.46919029407730
2 1 4		
3.50629783053394	10.6739641890506	231.756415050249
2.38789877391301	13.6673590924466	148.163889185864
2 2 3		
2.12383341617844	2.09610622027606	23.7246862784793
0.929943072656729	0.401190656024861	7.33017861737474
2 2 4		
2.10152459348052	9.94514189035131	240.500803642650
0.952091184635393	9.39249109396949	140.942518952472
2 3 4		
23.7104763915071	21.0346940586355	243.221560720708
7.34076531794134	-3.01793177595214	144.576874008222

The corresponding 'minimum' parameter file for a run including penalized estimation for a single tuning factor is:

	pool_min.par	
RUNOP --pool		1
ANAL MUV 4		2
VAR animal 4 NOS		3
VAR residual 4 NOS		4
POOL		5
MINPAR		6
SINGLE PartAll.dat		7
PSEUPED BON 100		8
DIRADD animal		9
PENALTY CORREL MAKETAR 2.5		10
END		11

Line 1 gives the `--pool` option, alternatively this could be specified on the command line. The first line in the `POOL` block (line 6) contains the `MINPAR` option, telling the program to expect minimal input, and line 7 directs the input to be read from the file `PartAll.dat`. Line 8 selects Bondari's design as pseudo pedigree structure with 100 families. With the `MINPAR` option, line 9 is required to instruct the program that the single random effect fitted is a direct, additive genetic effect. Finally, line 10 selects penalized pooling with shrinkage of the genetic towards the phenotypic covariance matrix using a tuning factor of 2.5, and line 11 closes the block.

The first part of `PoolEstimates.out` summarizes the numeric options, statistics on the input values, and gives the simple 'average' covariance matrices constructed from the part estimates together with their eigenvalues. With all possible, pairwise analyses between the 4 traits, there are 3 estimates for each variance but only one estimate for each covariance component.

	PoolEstimates.dat
===== Version 24-04-2012 ===== **KM** =====	
Program WOMBAT : Pooled estimates of covariance components	
=====	
Value for "small" = 0.00010000	
Convergence criterion = 0.00005000	
No. of traits = 4	
1 trait1	
2 trait2	
3 trait3	
4 trait4	
No. of part analyses = 6	
1 2-> 1 2 "PartAll.dat"	
2 2-> 1 3 "PartAll.dat"	
3 2-> 1 4 "PartAll.dat"	
4 2-> 2 3 "PartAll.dat"	
5 2-> 2 4 "PartAll.dat"	
6 2-> 3 4 "PartAll.dat"	
***** Means & ranges for residual covariances *****	
1 COVS Z 1 1 3 3.53132 3.50630 3.55743	
2 COVS Z 1 2 1 1.95049	

```

3 COVS Z 1 3      1      2.76836
4 COVS Z 1 4      1      10.6740
5 COVS Z 2 2      3      2.10481      2.08908      2.12383
6 COVS Z 2 3      1      2.09611
7 COVS Z 2 4      1      9.94514
8 COVS Z 3 3      3      23.6851      23.6201      23.7247
9 COVS Z 3 4      1      21.0347
10 COVS Z 4 4      3      238.493      231.756      243.222
Eigenvalues of averaged covariance matrix
241.466      21.8808      3.74837      0.719141

**** Means & ranges for RE 1 "animal" ****
11 COVS A 1 1      3      2.36108      2.32842      2.38790
12 COVS A 1 2      1      1.34230
13 COVS A 1 3      1      0.993394
14 COVS A 1 4      1      13.6674
15 COVS A 2 2      3      0.953153      0.929943      0.977426
16 COVS A 2 3      1      0.401191
17 COVS A 2 4      1      9.39249
18 COVS A 3 3      3      7.38004      7.33018      7.46919
19 COVS A 3 4      1      -3.01793
20 COVS A 4 4      3      144.561      140.943      148.164
Eigenvalues of averaged covariance matrix
146.540      7.63089      0.960639      0.124311

```

The second part gives details on the assumed pseudo pedigree and the results from unpenalized pooling. As the average matrices were within the parameter space and all analyses were weighted equally, not surprisingly, results differ little from the former.

```

PoolEstimates.dat continued

===== Pseudo pedigree structure =====
Bondari's design
No. of families      = 100
No. individuals/fam. = 8

Coefficients for RE 1 "animal"
1  1.0000
2  0.5000  1.0000
3  0.2500  0.2500  1.0000
4  0.2500  0.2500  0.5000  1.0000
5  0.5000  0.2500  0.1250  0.1250  1.0000
6  0.5000  0.2500  0.1250  0.1250  0.5000  1.0000
7  0.1250  0.1250  0.2500  0.5000  0.0625  0.0625  1.0000
8  0.1250  0.1250  0.2500  0.5000  0.0625  0.0625  0.5000  1.0000

=====
No. of parameters      = 20
**** Estimates of residual covariances ****
Covariance matrix
1  3.5450
2  1.9647      2.1023
3  2.7673      2.0763      23.672
4  11.131      9.8742      20.697      238.69
Eigenvalues of covariance matrix
Value      241.63      21.93      3.73      0.71
(%)      90.16      8.18      1.39      0.27
Trace      268.01
Matrix of correlations and variance ratios
1  0.6017
2  0.7197      0.6870
3  0.3021      0.2943      0.7621
4  0.3827      0.4408      0.2753      0.6233

**** Estimates for RE 1 "animal" ****
Covariance structure = NRM
Covariance matrix
1  2.3469
2  1.3219      0.95797
3  0.99423      0.42503      7.3899
4  13.338      9.5565      -2.9140      144.28
Eigenvalues of covariance matrix
Value      146.22      7.64      0.98      0.13
(%)      94.35      4.93      0.64      0.08
Trace      154.98
Matrix of correlations and variance ratios
1  0.3983
2  0.8816      0.3130
3  0.2387      0.1597      0.2379
4  0.7248      0.8129      -0.0892      0.3767

**** Estimates of phenotypic covariances ****
Covariance matrix
1  5.8919

```

```

 2    3.2865    3.0602
 3    3.7615    2.5013    31.062
 4    24.469    19.431    17.783    382.97
Eigenvalues of covariance matrix
Value    386.48    30.51    5.13    0.87
(%)      91.37    7.21    1.21    0.20
Trace    422.98
Correlation matrix
 1    1.0000
 2    0.7740    1.0000
 3    0.2780    0.2566    1.0000
 4    0.5151    0.5676    0.1630    1.0000
=====

```

Next, corresponding results when imposing a penalty are listed. For each covariance matrix, the change from the unpenalized, pooled estimates is summarized as the Frobenius norm (F-norm) of the matrix difference. If there were multiple tuning factors, such section would be given for each.

```

PoolEstimates.dat continued
**** Pooling with penalties ****
**** Type of penalty: "CORREL" ****

=====
===== Tuning factor = 2.50000 =====

**** Estimates of residual covariances ****
F-norm diff(unpenal) = 1.6007
Covariance matrix
 1    3.5590
 2    1.9826    2.1071
 3    2.8285    2.1021    23.670
 4    11.296    10.175    19.628    238.87
Eigenvalues of covariance matrix
Value    241.66    22.15    3.70    0.70
(%)      90.10    8.26    1.38    0.26
Trace    268.21
Matrix of correlations and variance ratios
 1    0.6046
 2    0.7240    0.6891
 3    0.3082    0.2977    0.7622
 4    0.3874    0.4535    0.2610    0.6242

**** Estimates for RE 1 "animal" ****
Covariance structure = NRM
F-norm diff(unpenal) = 2.1961
Covariance matrix
 1    2.3273
 2    1.2979    0.95079
 3    0.90708    0.38810    7.3847
 4    13.129    9.1775    -1.4612    143.82
Eigenvalues of covariance matrix
Value    145.64    7.58    1.12    0.15
(%)      94.27    4.91    0.73    0.10
Trace    154.49
Matrix of correlations and variance ratios
 1    0.3954
 2    0.8725    0.3109
 3    0.2188    0.1465    0.2378
 4    0.7176    0.7848    -0.0448    0.3758

**** Estimates of phenotypic covariances ****
F-norm diff(unpenal) = 0.62278
Covariance matrix
 1    5.8863
 2    3.2805    3.0579
 3    3.7356    2.4902    31.055
 4    24.426    19.352    18.167    382.70
Eigenvalues of covariance matrix
Value    386.23    30.45    5.14    0.87
(%)      91.37    7.20    1.22    0.21
Trace    422.70
Correlation matrix
 1    1.0000
 2    0.7732    1.0000
 3    0.2763    0.2555    1.0000
 4    0.5146    0.5657    0.1666    1.0000

```

Finally, for ease of comparison, results are listed side by side. For the combination of number of families, type of penalty and tuning factor, penalization did not affect

estimates substantially, but a slight reduction in the spread of the eigenvalues of the genetic covariance is notable.

```

PoolEstimates.dat continued
===== Estimates side by side =====
Tuning factor          0.00000      2.50000
Froben. Norm-Phen     0.00000      0.622778
Sum(Froben. Norm)     0.00000      3.79674

**** Estimates of residual covariances ****
 1 1 COVS Z 1 1  3.54499      3.55898
 1 2 COVS Z 1 2  1.96468      1.98260
 1 3 COVS Z 1 3  2.76727      2.82849
 1 4 COVS Z 1 4 11.1314      11.2964
 2 2 COVS Z 2 2  2.10228      2.10706
 2 3 COVS Z 2 3  2.07627      2.10211
 2 4 COVS Z 2 4  9.87423     10.1748
 3 3 COVS Z 3 3 23.6719     23.6703
 3 4 COVS Z 3 4 20.6967     19.6281
 4 4 COVS Z 4 4 238.690     238.873

**** Estimates for RE 1 "animal" ****
 1 1 COVS A 1 1  2.34691     2.32734
 1 2 COVS A 1 2  1.32185     1.29794
 1 3 COVS A 1 3  0.994227    0.907077
 1 4 COVS A 1 4 13.3376     13.1293
 2 2 COVS A 2 2  0.957967    0.950786
 2 3 COVS A 2 3  0.425032    0.388103
 2 4 COVS A 2 4  9.55651     9.17746
 3 3 COVS A 3 3  7.38991     7.38466
 3 4 COVS A 3 4 -2.91396    -1.46121
 4 4 COVS A 4 4 144.280     143.824

**** Estimates of phenotypic covariances ****
--- intermediate lines deleted ---

**** Eigenvalues for Residual ****
 1 E.value  241.633      241.657
 2 E.value  21.9283     22.1477
 3 E.value   3.73397     3.70123
 4 E.value   0.714643    0.702685

**** Eigenvalues for RE 1 "animal" ****
 1 E.value  146.218      145.638
 2 E.value   7.64321     7.57981
 3 E.value   0.984761     1.12056
 4 E.value   0.129158     0.148989
===== end of file =====26-04-2012=====15:31=====

```

4.2 More complicated example for 14 traits

Example 15 provides example runs from an analysis for 14 carcass and scan traits recorded on beef cattle. In addition to (penalized) pooling runs, corresponding runs for the methods proposed by Mäntysaari (1999) (run option `--it sum`) and Thompson et al. (2005) are shown.

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Appendix A Frequently asked questions

- Q:** I have a collection of estimates from various sources & data sets. Can I use WOMBAT to combine these?
A: Yes; the SINGLE input file and MINPAR options are provided to make ‘general’ use easier.
- Q:** My estimates are based on data sets of very different sizes. Can I do a weighted analysis?
A: Yes; each part analysis can be assigned a different weight. The degrees of freedom used in (Eq. 2) are then equal to the product of the weight, the number of families and the number of traits in the subset.
- Q:** I have existing covariance matrices which I would like to modify so that the smallest eigenvalues are a bit larger and, hopefully, my genetic evaluation runs converge faster. Can I do that with the --pool option?
A: Yes; WOMBAT can handle a single ‘part’ analysis with the number of traits in the ‘subset’ equal to the total number of traits. Beware though that setting the limit on the smallest eigenvalue to something quite different from zero can change matrices substantially - there is thus an upper limit of 1 on the value which can be set via the SMALL option. When attempting to regularize matrices, it might be best to combine a smaller limit on the minimum eigenvalues with a penalty.
- Q:** Can I get standard errors for the pooled estimates?
A: No; there is no theoretical basis to derive these.
- Q:** What value of tuning factor should I use?
A: The tuning factor specifies the emphasis given to the penalty relative to the ‘data’. Hence, the larger the assumed degrees of freedom (d_i in (Eq. 2) - which is proportional to the number of families in the pseudo pedigree structure, the number of traits and the weight given to a particular set of estimates), the less stringent the penalization for a given tuning factor. The general recommendation is to apply a ‘mild’ penalty - this may require trying a number of different values.
- Q:** Can I use --pool to modify a single matrix - and what do I need to watch out for?
A: The main value of likelihood based pooling lies in the opportunity to allow for repartitioning of the total variance into its components. However, yes this procedure can be applied to a single matrix. The parameter file should be set up calling this a residual covariance matrix - WOMBAT always expects to find one of these, but not necessarily any random effects and their covariance matrices. No pseudo pedigree structure is used in this case, and the parameter file thus should not include a PSEUPED statement. Similarly, the option MAKETAR for penalized estimation is not allowed - if penalized pooling with shrinkage towards a target matrix is required, this target needs to be specified explicitly. As canonical eigenvalues for a single matrix are not defined, the eigenvalues of the matrix are substituted for these if the option CANEIG is encountered.

Appendix B Technical details

B.1 Maximization of the likelihood

As the pseudo likelihood can be very flat, maximization is carried out very ‘heavy-handed’ using a Method of Scoring step followed by two derivative-free search steps, using methods of Powell (1965) and Nelder and Mead (1965), respectively. This is repeated until the change in log likelihood between rounds falls below a defined value (or if 50 rounds have been reached). For a stringent convergence criterion or a large number of traits, this can take quite some time.

B.2 Calculation of penalties

Penalties are summed over all sources of variation. For a penalty on the spread of canonical eigenvalues (CANEIG),

$$\mathcal{P} = \begin{cases} \sum_{x \neq E} \sum_{i=1}^q (\lambda_{xi} - \bar{\lambda}_x)^2 & \text{for ORG with } \bar{\lambda}_x = \sum_i \lambda_{xi} / q \\ \sum_{x \neq E} \sum_{i=1}^q \left(\log(\lambda_{xi}) - \overline{\log(\lambda_x)} \right)^2 + \left(\log(1 - \lambda_{xi}) - \overline{\log(1 - \lambda_x)} \right)^2 & \text{for LOG with } \overline{\log \lambda_x} = \sum_i \log(\lambda_{xi}) / q \end{cases}$$

and λ_{xi} the i -th canonical eigenvalue of $\hat{\Sigma}_p^{-1}\hat{\Sigma}_x$ and $\hat{\Sigma}_p = \sum_x \hat{\Sigma}_x$ the phenotypic covariance matrix. For this option, summation over x to obtain \mathcal{P} includes only the covariance components due to random effects fitted, i.e. excludes the residual covariance matrix.

Shrinking estimated covariance matrices towards a target matrix \mathbf{T} , the penalty is

$$\mathcal{P} = \sum_x \log |\hat{\Sigma}_x| + \text{tr}(\hat{\Sigma}_x^{-1}\mathbf{T})$$

with summation over x including all estimated covariance matrices. Invoking option MAKETAR sets \mathbf{T} to $\hat{\Sigma}_p$ from the unpenalized analysis. Similarly, the penalty for option CORREL is

$$\mathcal{P} = \sum_x \log |\hat{\mathbf{R}}_x| + \text{tr}(\hat{\mathbf{R}}_x^{-1}\mathbf{T})$$

with $\hat{\mathbf{R}}_x$ the correlation matrix corresponding to covariance matrix $\hat{\Sigma}_x$. Specifying MAKETAR for this case sets $\mathbf{T} = \hat{\mathbf{R}}_p$.

TO DO: add formulae for KANEIG and PACPAC