# movMF: An R Package for Fitting Mixtures of von Mises-Fisher Distributions

Kurt Hornik

Bettina Grün

WU Wirtschaftsuniversität Wien

Johannes Kepler Universität Linz

#### Abstract

Finite mixtures of von Mises-Fisher distributions allow to apply model-based clustering methods to data which is of standardized length, i.e., all data points lie on the unit sphere. The R package **movMF** contains functionality to draw samples from finite mixtures of von Mises-Fisher distributions and to fit these models using the expectation-maximization algorithm for maximum likelihood estimation. Special features are the possibility to use sparse matrix representations for the input data, different variants of the expectation-maximization algorithm, different methods for determing the concentration parameters in the M-step and to impose constraints on the concentration parameters over the groups.

In this paper we describe the main fitting function of the package and illustrate its application. We also discuss how several numerical issues are resolved which occur for estimating the concentration parameters and for determining the normalizing constant of the von Mises-Fisher distribution.

Keywords: EM algorithm, finite mixture, hypergeometric function  $_0F_1$ , modified Bessel function ratio, R, von Mises-Fisher distribution.

#### 1. Introduction

Finite mixture models allow to cluster observations by assuming that for each group a suitable parametric distribution can be specified and that the mixture distribution is derived by convex combination of the group distributions. An overview on the estimation of these models in a maximum likelihood as well as a Bayesian setting together with different applications of finite mixture models are presented in McLachlan and Peel (2000) and Frühwirth-Schnatter (2006).

If the support of the data is given by the unit sphere, a natural choice for the group distributions is the von Mises-Fisher (vMF) distribution. The special case where data is in  $\mathbb{R}^2$ , i.e., the observations lie on the unit circle, is referred to as von Mises distribution. The vMF distribution has concentric contour lines similar to the multivariate normal distribution with the variance-covariance matrix being a multiple of the identity matrix. In fact Mardia and Jupp (1999, p. 173) point out that if x follows a multivariate normal distribution with mean parameter  $\mu$  which has length one and variance-covariance matrix  $\kappa^{-1}I$ , then the conditional distribution of x given that it has length one is a vMF distribution with mean direction parameter  $\mu$  and concentration  $\kappa$ .

Finite mixtures of vMF distributions were introduced in Banerjee, Dhillon, Ghosh, and Sra (2005) to cluster data on the unit sphere. They propose to use the expectation-maximization (EM) algorithm for maximum likelihood (ML) estimation and present two variants which they

refer to as hard and soft clustering. The areas of applications of the presented examples include text mining where abstracts of scientific journals, news articles and newsgroup messages are categorized, and bioinformatics using a yeast gene expression data set. In these applications, available data is typically high-dimensional. The estimation methods employed need to take this into account. For finite mixtures of vMF distributions this is particularly relevant for the determination of the concentration parameters.

Tang, Chu, and Huang (2009) use finite mixtures of vMF distributions for speaker clustering. The data is pre-processed by fitting a Gaussian mixture model (GMM) to the utterances and stacking the mean vectors of the mixture components to form the GMM mean supervector which is then used as input for the mixture model aiming at clustering the speakers. They compare the performance of finite mixtures of Gaussian distributions with those of vMF distributions and conclude that for this application the latter outperform finite mixtures of multivariate normal distributions. Further possible areas of application for finite mixtures of vMFs are segmentation in high angular resolution diffusion imaging (McGraw, Vemuri, Yezierski, and Mareci 2006) and clustering treatment beam directions in external radiation therapy (Bangert, Hennig, and Oelfke 2010).

This paper is structured as follows. Section 2 introduces the vMF distribution and shows how to draw samples from this distribution and how to determine the ML estimates of its parameters. The extension to finite mixture models is covered in Section 3 including again the methods for drawing samples and determining ML estimates. The R (R Development Core Team 2012) package movMF is presented in Section 4 by describing the main fit function movMF() in detail. An illustrative application of the package using the abstracts from the talks at "useR! 2008", the 3rd international R user conference, in Dortmund, Germany, is given in Section 6. The paper concludes with a summary and an outlook on possible extensions.

### 2. The vMF distribution

A random unit length vector in  $\mathbb{R}^d$  has a *von Mises-Fisher* (or Langevin) distribution with parameter  $\theta \in \mathbb{R}^d$  if its density with respect to the uniform distribution on the unit sphere  $\mathbb{S}^{d-1} = \{x \in \mathbb{R}^d : ||x|| = 1\}$  is given by

$$f(x|\theta) = e^{\theta'x}/{}_{0}F_{1}(;d/2;||\theta||^{2}/4),$$

where

$$_{0}F_{1}(;\nu;z) = \sum_{n=0}^{\infty} \frac{\Gamma(\nu)}{\Gamma(\nu+n)} \frac{z^{n}}{n!}$$

is the confluent hypergeometric limit function and related to the modified Bessel function of the first kind  $I_{\nu}$  via

$$_{0}F_{1}(;\nu+1;z^{2}/4) = \frac{I_{\nu}(z)\Gamma(\nu+1)}{(z/2)^{\nu}}$$
 (1)

(e.g., Mardia and Jupp 1999, page 168).

The vMF distribution is commonly parametrized as  $\theta = \kappa \mu$ , where  $\kappa = \|\theta\|$  and  $\mu \in \mathbb{S}^{d-1}$  are the *concentration* and *mean direction* parameters, respectively (if  $\theta \neq 0$ ,  $\mu$  is uniquely determined as  $\theta/\|\theta\|$ ).

In what follows, it will be convenient to write

$$H_{\nu}(\kappa) = {}_{0}F_{1}(; \nu + 1; \kappa^{2}/4)$$

and

$$C_d(\kappa) = 1/{}_0F_1(; d/2; \kappa^2/4) = 1/H_{d/2-1}(\kappa)$$

so that the density is given by

$$f(x|\theta) = C_d(\|\theta\|)e^{\theta'x}.$$

#### 2.1. Simulating vMF distributions

The following algorithm provides a rejecting sampling scheme for drawing a sample from the vMF distribution with modal direction (0, ..., 0, 1) and concentration parameter  $\kappa \geq 0$  (see Algorithm VM\* in Wood 1994). The extension for simulating from the matrix Bingham vMF distribution is described in Hoff (2009) and also available in R through package **rstiefel** (Hoff 2012).

Step 1. Calculate b using

$$b = \frac{-2\kappa + \sqrt{4\kappa^2 + (d-1)^2}}{d-1}.$$

Put 
$$x_0 = (1-b)/(1+b)$$
 and  $c = \kappa x_0 + (d-1)\log(1-x_0^2)$ .

Step 2. Generate  $Z \sim \text{Beta}((d-1)/2, (d-1)/2)$  and  $U \sim \text{Unif}([0,1])$  and calculate

$$W = \frac{1 - (1 + b)Z}{1 - (1 - b)Z}.$$

Step 3. If

$$\kappa W + (d-1)\log(1-x_0W) - c < \log(U),$$

go to Step 2.

Step 4. Generate a uniform (d-1)-dimensional unit vector V and return

$$X = (\sqrt{1 - W^2}V', W)'.$$

The uniform (d-1)-dimensional unit vector V can be generated by simulating independent standard normal random variables and normalizing them (see for example Ulrich 1984). To get samples from a vMF distribution with arbitrary mean direction parameter  $\mu$ , X is multiplied from the left with a matrix where the first (d-1) columns consist of unitary basis vectors of the subspace orthogonal to  $\mu$  and the last column is equal to  $\mu$ .

#### 2.2. Estimating the parameters of the vMF distribution

Using the common parametrization by  $\kappa$  and  $\mu$ , the log-likelihood of a sample  $x_1, \ldots, x_n$  from the vMF distribution is given by

$$n\log(C_d(\kappa)) + \kappa \mu' r$$
,

where  $r = \sum_{i=1}^{n} x_i$  is the resultant vector (sum) of the  $x_i$ . The maximum likelihood estimates (MLEs) are obtained by solving the likelihood equations

$$\hat{\mu} = r/\|r\|, \qquad -\frac{C_d'(\hat{\kappa})}{C_d(\hat{\kappa})} = \|r\|/n.$$

Writing  $A_d(\kappa) = -C'_d(\kappa)/C_d(\kappa)$  for the logarithmic derivative of  $1/C_d(\kappa)$  and  $\rho = ||r||/n$  for average resultant length, the equation for the MLE of  $\kappa$  becomes

$$A_d(\hat{\kappa}) = \rho.$$

Using recursions for the modified Bessel functions (e.g., Watson 1995, page 71), one can establish that

$$A_d(\kappa) = -\frac{C'_d(\kappa)}{C_d(\kappa)} = \frac{I_{d/2}(\kappa)}{I_{d/2-1}(\kappa)}$$

so that using Equation 1,

$$A_d(\kappa) = \frac{(\kappa/2)^{\nu+1} H_{\nu+1}(\kappa) / \Gamma(\nu+2)}{(\kappa/2)^{\nu} H_{\nu}(\kappa) / \Gamma(\nu+1)} = \frac{\kappa}{d} \frac{H_{\nu+1}(\kappa)}{H_{\nu}(\kappa)}, \qquad \nu = d/2 - 1.$$

It can be shown (see for example Schou 1978) that  $A_d$  is a strictly increasing function which maps the interval  $[0, \infty)$  onto the interval [0, 1) and satisfies the Riccatti equation  $A'_d(\kappa) = 1 - A_d(\kappa)^2 - \frac{d-1}{\kappa} A_d(\kappa)$ . As  $A_d$  and hence its derivates can efficiently be computed using continued fractions (see Section 5 for details), the equation  $A_d(\hat{\kappa}) = \rho$  can efficiently be solved by standard iterative techniques provided that good starting approximations are employed.

Dhillon and Sra (2003) and subsequently Banerjee et al. (2005) suggest the approximation

$$\hat{\kappa} \approx \frac{\rho(d - \rho^2)}{1 - \rho^2} \tag{2}$$

obtained by truncating the Gauss continued fraction representation of  $A_d$  and adding a correction term "determined empirically". The former reference also suggests using this approximation as the starting point of a Newton-Raphson iteration, using the above expression for  $A'_d$ .

Tanabe, Fukumizu, Oba, Takenouchi, and Ishii (2007) show that

$$\hat{\kappa} = \frac{\rho(d-c)}{1-\rho^2}$$

for some suitable  $0 \le c \le 2$ . The approximation in Equation 2 corresponds to  $c \approx \rho^2$ . They also suggest to determine  $\hat{\kappa}$  via the fixed point iteration

$$\kappa_{t+1} = \kappa_t \rho / A_d(\kappa_t)$$

with a starting value in the solution range, e.g., using c = 1 or  $c = \rho^2$ .

Sra~(2012) suggests to use a "truncated Newton approximation" based on performing exactly two Newton iterations

$$\kappa_{t+1} = \kappa_t - \frac{A_d(\kappa_t) - \rho}{A_d'(\kappa_t)}$$

with  $\kappa_0$  determined via the  $c = \rho^2$  approximation.

Song, Liu, and Wang (2012) suggest to use a "truncated Halley approximation" based on performing exactly two Halley iterations

$$\kappa_{t+1} = \kappa_t - \frac{2(A_d(\kappa_t) - \rho)A'_d(\kappa_t)}{2A'_d(\kappa_t)^2 - (A_d(\kappa_t) - \rho)A''_d(\kappa_t)}$$

with  $\kappa_0$  determined via the  $c = \rho^2$  approximation and using that the second derivative can be given as a function of  $A_d(\kappa_t)$ , i.e.,

$$A_d''(\kappa_t) = 2A_d(\kappa_t)^3 + \frac{3(d-1)}{\kappa} A_d(\kappa_t)^2 + \frac{d^2 - d - 2\kappa^2}{\kappa^2} A_d(\kappa_t) - \frac{d-1}{\kappa}.$$

The results in Hornik and Grün (2012) yield the substantially improved bounds for  $0 \le \rho < 1$ 

$$\max\left(F_{d/2-1,d/2+1}(\rho), F_{(d-1)/2,\sqrt{d^2-1}/4}(\rho)\right) \le \hat{\kappa} \le F_{(d-1)/2,(d+1)/2}(\rho),\tag{3}$$

where

$$F_{\alpha,\beta}(\rho) = \frac{\rho}{1 - \rho^2} \left( \alpha + \sqrt{\rho^2 \alpha^2 + (1 - \rho^2)\beta^2} \right).$$

The difference between the upper and lower bound is at most  $3\rho/2$  for all  $0 \le \rho < 1$ , and the difference between the lower bound and  $\hat{\kappa}$  tends to 0 as  $\rho \to 1-$ .

Clearly, the availability of such bounds makes it possible to determine  $\hat{\kappa}$  using numerical methods which are able to detect the root of a function inside a specified interval (such as uniroot() in R).

#### 3. Finite mixtures of vMF distributions

The mixture model with K components which correspond to K groups is given by

$$h(x|\Theta) = \sum_{k=1}^{K} \pi_k f(x|\theta_k),$$

where  $h(\cdot|\cdot)$  is the mixture density,  $\Theta$  the vector with all  $\pi$  and  $\theta$  parameters, and  $f(y|\theta_k)$  the density of the vMF distribution with parameter  $\theta_k$ . Furthermore the component weights  $\pi_k$  are non-negative for all k and sum to one.

#### 3.1. Simulating mixtures of vMF distributions

This is straightforwardly achieved by first sampling class ids  $z \in \{1, ..., K\}$  with the mixture class probabilities  $\pi_1, ..., \pi_K$ , and then sampling the data from the respective vMF distributions with parameter  $\theta_k$ .

### 3.2. Estimating the parameters of mixtures of vMF distributions

EM algorithms for ML estimation of the parameters of mixtures of vMF distributions are given in Dhillon and Sra (2003) and Banerjee *et al.* (2005). The EM algorithm exploits that fact that the complete-data log-likelihood where the component memberships of the observations are known is easier to maximize than the observed-data log-likelihood.

The EM algorithm for fitting mixtures of vMF distributions consists of the following steps:

- 1. Initialization: Either of the following two:
  - (a) Assign values to  $\pi_k$  and  $\theta_k$  for k = 1, ..., K, where  $\pi_k > 0$  and  $\sum_{k=1}^K \pi_k = 1$  and  $\theta_k \neq \theta_l$  for all  $k \neq l$  and k, l = 1, ..., K. Start the EM algorithm with an E-step.
  - (b) Assign (probabilities of) component memberships to each of the n observations. E.g., the output from spherical k-means (Hornik, Feinerer, Kober, and Buchta 2012) can be used.

Start the EM algorithm with an M-step.

2. Repeat the following steps until the maximum number of iterations is reached or the converence criterion is met.

**E-step:** Because the complete-data log-likelihood is linear in the missing data which correspond to the component memberships, the E-step only consists of calculating the a-posteriori probabilities, the probabilities of belonging to an component conditional on the observed values, using

$$p_{ik} := p(k|x_i, \Theta) \propto \pi_k f(x_i|\theta_k).$$

**M-step:** Maximize the expected complete-data log-likelihood by determining separately for each k:

$$\hat{\pi}_{k} = \frac{1}{n} \sum_{i=1}^{n} p(k|x_{i}, \Theta),$$

$$\hat{\mu}_{k} = \frac{\sum_{i=1}^{n} p(k|x_{i}, \Theta)x_{i}}{\|\sum_{i=1}^{n} p(k|x_{i}, \Theta)x_{i}\|}, \qquad -\frac{C'_{d}(\hat{\kappa}_{k})}{C_{d}(\hat{\kappa}_{k})} = \frac{\|\sum_{i=1}^{n} p(k|x_{i}, \Theta)x_{i}\|}{\sum_{i=1}^{n} p(k|x_{i}, \Theta)}.$$

Convergence check: Assess convergence by checking

- (a) if the relative absolute change in the log-likelihood values is smaller than a threshold  $\epsilon_1$  and / or
- (b) if the relative absolute change in parameters is smaller than a threshold  $\epsilon_2$ .

If converged, stop the algorithm.

In the following we will only use condition (a) to check for convergence.

This corresponds to the soft-movMF algorithm on page 1357 in Banerjee *et al.* (2005). In addition they propose the hard-movMF algorithm on page 1358. The algorithm above can be modified to the hard-movMF algorithm by adding a hardening step between E- and M-step:

**H-step:** Replace the a-posteriori probabilities by assigning each observation with probability 1 to the component where its a-posteriori probability is maximum.

$$p(k|x_i, \Theta) = \begin{cases} 1, & \text{if } k = \operatorname{argmax}_h p_{ih} \\ 0, & \text{otherwise} \end{cases}$$

 $\hat{\kappa}_k$  in the M-step can be determined via the approximation of Equation (2), or one of the improved methods discussed in Section 2.2.

The algorithm above determines the parameter estimates if the concentration parameters are allowed to vary freely over components. An alternative model specification could be to impose the restriction that the concentration parameters are the same for each component. This has the advantage that the groups will be of comparable compactness and that spurious solutions containing small components with very large concentration parameters are eliminated. In the following we derive how the M-step needs to be modified if the concentration parameters are restricted to be the same over components.

From Appendix A.2 of Banerjee et al. (2005) the optimal unconstrained  $\kappa_k$  can be obtained by solving

$$A_d(\kappa_k) = \frac{\|\sum_i p(k|x_i, \Theta)x_i\|}{\sum_i p(k|x_i, \Theta)}.$$

If the  $\kappa_k$  are constrained to be equal (but are not given), the optimal common  $\kappa$  can be obtained as follows. Using Equation A.12 in Banerjee *et al.* (2005), the modified Lagrangian becomes

$$\sum_{k} \left( \sum_{i} \log(C_d(\kappa)) p(k|x_i, \Theta) + \sum_{i} \kappa \mu'_k x_i p(k|x_i, \Theta) \right) + \lambda_k (1 - \mu'_k \mu_k)$$

$$= \log(C_d(\kappa)) \sum_{k,i} p(k|x_i, \Theta) + \kappa \sum_{k,i} \mu'_k x_i p(k|x_i, \Theta) + \lambda_k (1 - \mu'_k \mu_k)$$

$$= n \log(C_d(\kappa)) + \kappa \sum_{k,i} \mu'_k x_i p(k|x_i, \Theta) + \lambda_k (1 - \mu'_k \mu_k).$$

Setting partials with respect to  $\mu_k$  and  $\lambda_k$  to zero as in the reference gives (again)

$$\mu_k = \frac{\sum_i x_i p(k|x_i, \Theta)}{\|\sum_i x_i p(k|x_i, \Theta)\|}$$

and with these  $\mu_k$  we obtain for  $\kappa$  that

$$0 = -A_d(\kappa)n + \sum_{k,i} \mu'_k x_i p(k|x_i, \Theta) = -A_d(\kappa)n + \sum_k \left\| \sum_i x_i p(k|x_i, \Theta) \right\|,$$

i.e.,  $\kappa$  needs to solve the equation

$$A_d(\kappa) = \frac{1}{n} \sum_{k} \left\| \sum_{i} x_i p(k|x_i, \Theta) \right\|.$$

### 4. Main fitting function movMF()

The main function in package **movMF** for fitting mixtures of vMF distributions is **movMF()**, with synopsis

```
R> movMF(x, k, control = list(), ...)
```

The arguments for this function are as follows.

- x: A numeric data matrix, with rows corresponding to observations. If necessary the data is standardized to unit row lengths. Furthermore, the matrix can be either stored as a dense matrix, a simple triplet matrix (defined in package slam), or a general sparse triplet matrix of class "dgtMatrix" (from package Matrix).
- k: An integer indicating the number of components.

control: A list of control parameters consisting of

E: Specifies the variant of the EM algorithm used with possible values "softmax" (default), "hardmax" (Classification EM), and "stochmax" (Stochastic EM).

kappa: This argument allows to specify how to determine the concentration parameters.

- If numbers are given, the concentration parameters are assumed to be fixed and are not estimated in the EM algorithm.
- The method for solving for the concentration parameters can be specified by one of "Banerjee\_et\_al\_2005", "Tanabe\_et\_al\_2007", "Sra\_2012", "Song\_et\_al\_2012", "uniroot", "Newton" (default).

  Here, the first four correspond to the approximation, the fixed point iteration and the truncated Newton and Halley approximation, respectively, of Banerjee et al. (2005), Tanabe et al. (2007), Sra (2012) and Song et al. (2012) described in Section 2.2. Method "uniroot" uses a straightforward call to uniroot() with initials bounds from Equation 3. Finally, method "Newton" performs a full Newton-Raphson iteration with the lower bound from Equation 3 as starting value.
- For common concentration parameters a list with elements common = TRUE and a character string giving the estimation method needs to be provided.
- converge: Logical indicating if convergence of the algorithm should be checked and if in this case the algorithm should be stopped before the maximum number of iterations is reached. For E equal to "softmax" this argument is set to FALSE.
- maxiter: Integer indicating the maximum number of iterations of the EM algorithm. (Default: 100.)
- reltol: If the relative change in the log-likelihood falls below this threshold the EM algorithm is stopped if converge is TRUE. (Default: sqrt(.Machine\$double.eps).)
- **verbose:** Logical indicating if information on the progress of the fitting process shall be printed during the estimation.

ids: Indicates either the class memberships of the observations or if equal to TRUE the class memberships are obtained from the attributes of the data. In this way the class memberships are for example stored if data is generated using function rmovMF(). In this case the EM algorithm is stopped after one iteration, i.e., the parameter estimates are determined conditional on the known true class memberships.

start: Allows to specify the starting values used in the EM algorithm. It can either be a list of matrices where each matrix contains in each row the a-posteriori probabilities of the observations or a list of vectors containing component assignments for the observations. Alternatively it can be a character vector with entries "i", "p", "S" or "s". The length of the vector specifies how many different initializations are made. "i" indicates to randomly assign component memberships to the observations. The latter three draw observations as prototypes and determine a-posteriori probabilities by taking the implied cosine dissimilarities between observations and prototypes. "p" randomly picks observations as prototypes, "S" takes the first prototype to minimize total cosine dissimilarity to the observations, and then successively picks observations farthest away from the already picked prototypes. For "s" one takes a randomly chosen observation as the first prototype, and then proceeds as for "S".

nruns: An integer indicating the number of repeated runs of the EM algorithm with random initialization. This argument is ignored if either ids or start are specified.

minalpha: Components with size below the threshold indicated by minalpha are omitted during the estimation with the EM algorithm. This avoids estimation problems in the M-step if only very few observations are assigned to one component. The disadvantage is that the initial number of components is not necessarily equal to the number of components of the returned model.

The object returned by movMF() is an S3 class called "movMF" with methods print(), coef(), logLik() and predict() (obtains either the component assignments or the matrix of a-posteriori probabilities).

Additional functionality available in the package includes rmovMF() for drawing from a mixture of vMF distributions and dmovMF() for evaluating the mixture density.

### 5. Numerical Issues

As previously shown, computing log-likelihoods for (mixtures of) vMF distributions on  $\mathbb{R}^d$  requires the computation of  $\log(H_{d/2-1})$ , where

$$H_{\nu}(\kappa) = {}_{0}F_{1}(; \nu + 1; \kappa^{2}/4) = \frac{\Gamma(\nu + 1)}{(\kappa/2)^{\nu}}I_{\nu}(\kappa),$$

and ML estimation of concentration parameters amounts to solving equations of the form  $A_d(\kappa) = \rho$ , where

$$A_d(\kappa) = \frac{I_{d/2}(\kappa)}{I_{d/2-1}(\kappa)} = \frac{\kappa}{d} \frac{H_{d/2}(\kappa)}{H_{d/2-1}(\kappa)}$$

is the logarithmic derivative of  $H_{d/2-1}$ .

As  $H_{\nu}(\kappa) \to \infty$  as  $\kappa \to \infty$  (in fact, quite rapidly, see below), it clearly is a bad idea to try to compute  $\log(H)$  as the logarithm of H, or A as the ratio of H functions. Similar considerations apply for using logarithms or ratios of incomplete modified Bessel functions I. One might wonder whether one could successfully take advantage of the fact that the Bessel functions provided by R are based on the **SPECFUN** package of Cody (1993) and hence also provide the exponentially scaled modified Bessel function  $e^{-\kappa}I_{\nu}(\kappa)$  (the scaling is motivated by the asymptotic expansion  $I_{\nu}(\kappa) \sim e^{\kappa}(2\pi\kappa)^{-1/2}\sum_{m}\alpha_{m}(\nu)/\kappa^{m}$  for  $\kappa \to \infty$ ). However, exponential scaling does not help in situations where  $1 \ll \kappa \ll \nu$ : e.g., for  $\kappa = 6000$  and  $\nu = 10000$ , H overflows whereas I underflows (even though R gives Inf and 0 for the cases without and with exponential scaling, respectively).

One can use the Gauss continued fraction

$$\frac{I_{\nu}(z)}{I_{\nu-1}(z)} = \frac{1}{2\nu/z + 2(\nu+1)/z + 2(\nu+2)/z + \cdots}$$

for the ratio of modified Bessel functions (http://dlmf.nist.gov/10.33.E1) to compute A as

$$A_d(\kappa) = \frac{I_{d/2}(\kappa)}{I_{d/2-1}(\kappa)} = \frac{1}{d/\kappa + \frac{1}{(d+2)/\kappa + \frac{1}{(d+4)/\kappa + \cdots}}} \cdots$$

(Equation 4.3 in Banerjee et al. 2005), using, e.g., Steed's method (e.g., http://dlmf.nist.gov/3.10) for evaluation. However, as pointed out by Gautschi and Slavik (1978) and Tretter and Walster (1980) (and, quite recently, re-iterated by Song et al. 2012), the Perron continued fraction

is numerically more stable (as computing it by forward recursion only accumulates positive terms, whereas for the Gauss continued fraction the terms alternate in sign), and converges substiantally faster for positive  $z \gg \nu$ . Hence, by default we compute A via the Perron continued fraction (with implementation based on Equation 3.3' in Gautschi and Slavik 1978), and additionally provide computation via the Gauss continued fraction (with implementation based on Equation 3.2' in Gautschi and Slavik 1978) and using exponentiation of  $\log(H)$  differences as alternatives.

For  $\kappa$  close to zero it is better (and necessary for  $\kappa = 0$ ) to use the approximation

$$A_d(\kappa) = \frac{1}{d}\kappa - \frac{1}{d^2(d+2)}\kappa^3 + O(\kappa^5), \qquad \kappa \to 0$$

(Schou 1978, Equation 5). The  $O(\kappa)^5$  can be made more precise by using the series representation  $I_{\nu}(z) = \sum_{n=0}^{\infty} (z/2)^{2n+\nu}/(n!\Gamma(n+\nu+1))$  so that for  $\kappa \to 0$ ,

$$A_{d}(\kappa) = \frac{(\kappa/2)^{d/2} \left(\frac{1}{\Gamma(d/2+1)} + \frac{\kappa^{2}/4}{\Gamma(d/2+2)} + \frac{\kappa^{4}/32}{\Gamma(d/2+3)} + O(\kappa^{6})\right)}{(\kappa/2)^{d/2-1} \left(\frac{1}{\Gamma(d/2)} + \frac{\kappa^{2}/4}{\Gamma(d/2+1)} + \frac{\kappa^{2}/32}{\Gamma(d/2+2)} + O(\kappa^{6})\right)}$$

$$= \frac{\kappa}{2} \frac{\frac{2}{d} + \frac{\kappa^{2}}{d(d+2)} + \frac{\kappa^{4}}{4d(d+2)(d+4)} + O(\kappa^{6})}{1 + \frac{\kappa^{2}}{2d} + \frac{\kappa^{4}}{8d(d+2)} + O(\kappa^{6})}$$

from which the coefficient of  $\kappa^5$  can straightforwardly be determined as

$$\frac{1}{2} \left( \frac{1}{4d(d+2)(d+4)} - \frac{1}{2d^2(d+2)} - \frac{2}{8d^2(d+2)} + \frac{2}{4d^3} \right) = \frac{2}{d^3(d+2)(d+4)}$$

so that

$$A_d(\kappa) = \frac{1}{d}\kappa - \frac{1}{d^2(d+2)}\kappa^3 + \frac{2}{d^3(d+2)(d+4)}\kappa^5 + O(\kappa^7), \qquad \kappa \to 0.$$

From this approximations for A' and A'' can be obtained by term-wise differentiation and also used for  $\kappa$  close to 0.

Write

$$R_{\nu}(\kappa) = \frac{I_{\nu+1}(\kappa)}{I_{\nu}(\kappa)}$$

for the Bessel function ratio (so that  $A_d = R_{d/2-1}$ ) and

$$G_{\alpha,\beta}(\kappa) = \frac{\kappa}{\alpha + \sqrt{\kappa^2 + \beta^2}}.$$

Amos (1974, Equations 11 and 16) shows that for all non-negative  $\kappa$  and  $\nu$ ,  $G_{\nu+1/2,\nu+3/2}(\kappa) \leq R_{\nu}(\kappa) \leq G_{\nu,\nu+2}(\kappa)$ . With  $\beta_{SS}(\nu) = \sqrt{(\nu+1/2)(\nu+3/2)}$ , Theorem 2 of Simpson and Spector (1984) implies the upper bound  $R_{\nu}(\kappa) \leq G_{\nu+1/2,\beta_{SS}(\nu)}(\kappa)$  for all non-negative  $\kappa$  and  $\nu$ . We thus have

$$G_{\nu+1/2,\nu+3/2}(\kappa) \le R_{\nu}(\kappa) \le \min \left( G_{\nu,\nu+2}(\kappa), G_{\nu+1/2,\beta_{SS}(\nu)}(\kappa) \right)$$
 (4)

(from which the bounds of Equation 3 are obtained by inversion). In addition, using results in Schou (1978, Equations 5 and 6), one can show that  $G_{\nu,\nu+2}$  and  $G_{\nu+1/2,\beta_{SS}(\nu)}$  are second order exact approximations for  $\kappa \to 0$  and  $\kappa \to \infty$ , respectively.

As  $\log(H_{\nu})' = R_{\nu}$  (and  $H_{\nu}(0) = 1$ ), integration gives

$$\log(H_{\nu}(\kappa)) = \int_{0}^{\kappa} R_{\nu}(t) dt.$$

Thus, the bounds for the Bessel function ratio  $R_{\nu}$  can be used to obtain bounds for  $H_{\nu}$ . Writing

$$S_{\alpha,\beta}(\kappa) = \sqrt{\kappa^2 + \beta^2} - \alpha \log(\alpha + \sqrt{\kappa^2 + \beta^2}) - \beta + \alpha \log(\alpha + \beta),$$

it is easily verified that  $S'_{\alpha,\beta} = G_{\alpha,\beta}$  and  $S_{\alpha,\beta}(0) = 0$ . Using the Amos-type bounds from Equation 4, we thus obtain that for  $\nu \geq 0$  and  $\kappa \geq 0$ ,

$$S_{\nu+1/2,\nu+3/2}(\kappa) \le \log(H_{\nu}(\kappa)) \le \min(S_{\nu,\nu+2}(\kappa), S_{\nu+1/2,\beta_{SS}(\nu)}(\kappa)).$$
 (5)

Where a single approximating value is sought, we prefer to use the upper bound which is based on the combination of upper Amos-type bounds which are second order exact at zero and infinity. Again, the bounds in Equation 5 are surprisingly tight.

Result. Let

$$s(\nu) = (\nu + 3/2) - \beta_{SS}(\nu) - (\nu + 1/2) \log \frac{2(\nu + 1)}{\nu + 1/2 + \beta_{SS}(\nu)},$$

with  $\nu \geq 0$ .

Then s is non-increasing on  $[0, \infty)$  with  $s(0) = (3 - \sqrt{3} + \log((1 + \sqrt{3})/4))/2 = 0.4433537$  and  $\lim_{\nu \to \infty} s(\nu) = 1/4$ . For  $\nu_0 \ge 0$ ,

$$\sup_{\kappa \ge 0, \nu \ge \nu_0} (S_{\nu+1/2, \beta_{SS}(\nu)}(\kappa) - S_{\nu+1/2, \nu+3/2}(\kappa)) = s(\nu_0).$$

For  $\beta_{SS}(\nu) \le \beta \le \nu + 3/2$ ,

$$\sup_{\kappa \ge 0} |\log(H_{\nu}(\kappa)) - S_{\nu+1/2,\beta}(\kappa)| \le s(\nu).$$

*Proof.* For simplicity, write  $\alpha = \nu + 1/2$ ,  $\beta_L = \nu + 3/2$  and  $\beta_U = \beta_{SS}(\nu)$ , omitting the dependence on  $\nu$ . We have  $\beta_U \leq \beta_L$  and  $G_{\alpha,\beta_U} \geq G_{\alpha,\beta_L}$ . Hence,

$$S_{\alpha,\beta_U}(\kappa) - S_{\alpha,\beta_L}(\kappa) = \int_0^{\kappa} (G_{\alpha,\beta_U}(t) - G_{\alpha,\beta_L}(t)) dt$$

is non-decreasing in  $\kappa$ , and attains its supremum for  $\kappa \to \infty$ . Now,

$$S_{\alpha,\beta_U}(\kappa) - S_{\alpha,\beta_L}(\kappa) = \sqrt{\kappa^2 + \beta_U^2} - \sqrt{\kappa^2 + \beta_L^2} - \alpha \log \left( \frac{\alpha + \sqrt{\kappa^2 + \beta_U^2}}{\alpha + \sqrt{\kappa^2 + \beta_L^2}} \right) + (\beta_L - \beta_U) - \alpha \log \frac{\alpha + \beta_L}{\alpha + \beta_U}.$$

As

$$\sqrt{\kappa^2 + \beta_U^2} - \sqrt{\kappa^2 + \beta_L^2} = \frac{(\kappa^2 + \beta_U^2) - (\kappa^2 + \beta_L^2)}{\sqrt{\kappa^2 + \beta_U^2} + \sqrt{\kappa^2 + \beta_L^2}} = \frac{\beta_U^2 - \beta_L^2}{\sqrt{\kappa^2 + \beta_U^2} + \sqrt{\kappa^2 + \beta_L^2}},$$

we have

$$\lim_{\kappa \to \infty} (S_{\alpha, \beta_U}(\kappa) - S_{\alpha, \beta_L}(\kappa)) = (\beta_L - \beta_U) - \alpha \log \frac{\alpha + \beta_L}{\alpha + \beta_U} = s(\nu).$$

The value of s(0) is obtained by insertion, and  $\lim_{\nu\to\infty} s(\nu)$  can be obtained as follows. We have

$$\beta_L - \beta_U = \sqrt{\alpha + 1}(\sqrt{\alpha + 1} - \sqrt{\alpha}) = \frac{\sqrt{\alpha + 1}}{\sqrt{\alpha + 1} + \sqrt{\alpha}} \to 1/2$$

as  $\nu \to \infty$ , and

$$\begin{split} \frac{\alpha + \beta_U}{\alpha + \beta_L} &= \frac{\alpha + \sqrt{\alpha(\alpha + 1)}}{2\alpha + 1} \\ &= \frac{(1 + \sqrt{1 + 1/\alpha})/2}{1 + 1/(2\alpha)} \\ &= \frac{1}{2} \left( 1 + 1 + \frac{1}{2\alpha} + O(\alpha^{-2}) \right) \left( 1 - \frac{1}{2\alpha} + O(\alpha^{-2}) \right) \\ &= 1 - \frac{1}{4\alpha} + O(\alpha^{-2}) \end{split}$$

so that

$$\alpha \log \frac{\alpha + \beta_U}{\alpha + \beta_L} = \alpha \left( -\frac{1}{4\alpha} + O(\alpha^{-2}) \right) = -\frac{1}{4} + O(\alpha^{-1}) \to -1/4$$

as  $\nu \to \infty$ . Hence,  $\lim_{\nu \to \infty} s(\nu) = 1/4$ .

To show that s is non-increasing, note that we have

$$\alpha + \beta_L = 2\alpha + 1,$$
  $\frac{d\alpha}{d\nu} = \frac{d\beta_L}{d\nu} = 1,$   $\frac{d\beta_U}{d\nu} = \frac{2\alpha + 1}{2\beta_U}$ 

and hence

$$\frac{ds}{d\nu} = 1 - \frac{2\alpha + 1}{2\beta_U} - \log \frac{2\alpha + 1}{\alpha + \beta_U} - \alpha \left( \frac{2}{2\alpha + 1} - \frac{1}{\alpha + \beta_U} \left( 1 + \frac{2\alpha + 1}{2\beta_U} \right) \right).$$

For non-negative t and  $\tau$ ,

$$\log \frac{t+\tau}{t} = \int_0^\tau \frac{1}{t+s} \, ds \ge \frac{1}{t+\tau} \int_0^\tau ds = \frac{\tau}{t+\tau},$$

Hence, with  $t = \alpha + \beta_U$  and  $\tau = \beta_L - \beta_U$ ,

$$\log \frac{2\alpha + 1}{\alpha + \beta_U} \ge \frac{\beta_L - \beta_U}{2\alpha + 1}$$

and

$$\frac{ds}{d\nu} \leq 1 - \frac{2\alpha + 1}{2\beta_U} + \frac{\beta_U - \beta_L}{2\alpha + 1} + \frac{\alpha}{\alpha + \beta_U} \left( 1 + \frac{2\alpha + 1}{2\beta_U} \right) - \frac{2\alpha}{2\alpha + 1} 
= \frac{2\alpha + 1}{2\beta_U} \left( \frac{\alpha}{\alpha + \beta_U} - 1 \right) + \frac{\alpha}{\alpha + \beta_U} + \frac{2\alpha + 1 + \beta_U - \beta_L - 2\alpha}{2\alpha + 1} 
= -\frac{2\alpha + 1}{2(\alpha + \beta_U)} + \frac{\alpha}{\alpha + \beta_U} + \frac{\beta_U - \alpha}{2\alpha + 1} 
= -\frac{1}{2(\alpha + \beta_U)} + \frac{\beta_U - \alpha}{2\alpha + 1} 
= \frac{-(2\alpha + 1) + 2(\beta_U^2 - \alpha^2)}{2(\alpha + \beta_U)(2\alpha + 1)} 
= \frac{-1}{2(\alpha + \beta_U)(2\alpha + 1)} 
\leq 0,$$

establishing that s is non-increasing.

Finally, for  $\beta_{SS}(\nu) \leq \beta \leq \nu + 3/2$ , both  $\log(H_{\nu})$  and  $S_{\nu+1/2,\beta}$  are bounded below by  $S_{\nu+1/2,\nu+3/2}$  and above by  $S_{\nu+1/2,\beta_{SS}(\nu)}$ , so that  $|\log(H_{\nu}) - S_{\nu+1/2,\beta}| \leq (S_{\nu+1/2,\beta_{SS}(\nu)} - S_{\nu+1/2,\nu+3/2}) \leq s(\nu)$ , completing the proof.

These results can be used to derive the following approach to computing  $\log(H_{\nu})$ . Choose a threshold  $\theta$  such that  $e^{\theta}$  does not overflow and  $e^{-\theta}$  does not underflow. Using IEEE 754 double precision floating point computations, we can take  $\theta = 700$ . Compute some convex combination of the lower and upper bounds in Equation 5 (e.g., simply take the upper bound) or use  $S_{\nu+1/2,\beta}(\kappa)$  with some  $\beta_{SS}(\nu) \leq \beta \leq \nu + 3/2$  (e.g.,  $\beta = \nu + 1$ ) to obtain an approximation  $L_{\nu}(\kappa)$  to  $\log(H_{\nu}(\kappa))$  which we know has approximation error at

most s(0) < 1/2. If  $L_{\nu}(\kappa) \le \theta - 1/2$  (so that  $\log(H_{\nu}(\kappa)) \le \theta$ ), compute  $H_{\nu}(\kappa)$  by its series expansion, and take the logarithm of this. If  $\theta - 1/2 < L_{\nu}(\kappa) \le 2\theta - 1$  so that

$$|\log(H_{\nu}(\kappa)) - L_{\nu}(\kappa)/2| \le |\log(H_{\nu}(\kappa)) - L_{\nu}(\kappa)| + L_{\nu}(\kappa)/2 \le \theta,$$

and hence  $e^{-L_{\nu}(\kappa)/2}H_{\nu}(\kappa)$  does not over- or underflow, write

$$H_{\nu}(\kappa) = e^{L_{\nu}(\kappa)/2} e^{-L_{\nu}(\kappa)/2} H_{\nu}(\kappa) = e^{L_{\nu}(\kappa)/2} \sum_{m=0}^{\infty} \frac{e^{-L_{\nu}(\kappa)/2} \Gamma(\nu+1)}{\Gamma(\nu+1+m)} \frac{(\kappa^2/4)^m}{m!},$$

and compute  $\log(H_{\nu}(\kappa))$  as  $L_{\nu}(\kappa)/2$  plus the logarithm of the sum of the scaled series. Otherwise, use the approximation  $L_{\nu}(\kappa)$  (note that with  $\theta = 700$ , this has a relative approximation error of about at most  $1/2 \cdot 1/1400 \le 0.0004$ ). This is the approach for computing  $\log(H_{\nu})$  currently implemented in package **movMF**, using the upper bound in Equation 5 as approximation  $L_{\nu}$  to  $\log(H_{\nu})$ .

Alternatively, one can use the bounds to obtain refined strategies of computing log(H) using available codes for modified Bessel functions. We have

$$\log(H_{\nu}(\kappa)) = \log(I_{\nu}(\kappa)) - \nu \log(\kappa/2) + \log(\Gamma(\nu+1))$$

and hence, using Stirling's approximation  $\log(\Gamma(z)) \approx (z - 1/2) \log(z) - z + \log(2\pi)/2$  and  $L_{\nu} = S_{\nu+1/2,\nu+1}$  for notational convenience, gives

$$\begin{split} \log(I_{\nu}(\kappa)) &= \log(H_{\nu}(\kappa)) + \nu \log(\kappa/2) - \log(\Gamma(\nu+1)) \\ &\approx L_{\nu}(\kappa) + \nu \log(\kappa/2) - (\nu+1/2) \log(\nu+1) + (\nu+1) - \frac{\log(2\pi)}{2} \\ &= \sqrt{\kappa^2 + (\nu+1)^2} + (\nu+1/2) \log \frac{\kappa}{\nu + 1/2 + \sqrt{\kappa^2 + (\nu+1)^2}} \\ &- \frac{\log(\kappa/2)}{2} + (\nu+1/2) \log \frac{2\nu + 3/2}{2(\nu+1)} - \frac{\log(2\pi)}{2}. \end{split}$$

This implies

$$\log(I_{\nu}(\kappa)) = \sqrt{\kappa^2 + (\nu+1)^2} + (\nu+1/2)\log\frac{\kappa}{\nu + 1/2 + \sqrt{\kappa^2 + (\nu+1)^2}} - \frac{\log(\kappa)}{2} + O(1),$$

where the O(1) can be made more precise, giving a first-order variant of the large  $\nu$  uniform asymptotic approximation for  $I_{\nu}$  given by Olver (1954). (For  $\nu \to \infty$ , the error in the approximation for  $\log(\Gamma)$  tends to zero, and as  $(\nu+1/2)\log((2\nu+3/2)/(2\nu+2)) \to -1/4$ , the O(1) becomes  $-\log(\pi)/2 - 1/4 + o(1)$  (modulo the error in the approximation of  $\log(H_{\nu}(\kappa))$  by  $L_{\nu}(\kappa)$  which is at most 1/4 + o(1)).

From the above, we can see that when  $\kappa \gg \nu$ ,  $I_{\nu}(\kappa)$  overflows quite rapidly; on the other hand, for  $\kappa = o(\nu)$  and  $\nu \to \infty$ ,  $I_{\nu}(\kappa)$  underflows quite rapidly. For computing logarithms, overflow can be avoided by employing the exponentially scaled modified Bessel function  $e^{-\kappa}I_{\nu}(\kappa)$  (as commonly available in codes for computing Bessel fuctions, such as the **SPECFUN** package Cody 1993, used by R) and computing  $\log(I_{\nu}(\kappa)) = \kappa + \log(e^{-\kappa}I_{\nu}(\kappa))$ . However, this clearly does not help avoiding underflow.

This motivates the following strategy for computing  $\log(I_{\nu}(\kappa))$  for a wide range of values. Start by computing a quick approximation  $T_{\nu}(\kappa)$  to  $\log(I_{\nu})$ , either using the above  $L_{\nu}(\kappa) + \nu \log(\kappa/2) - \log(\Gamma(\nu+1))$  or the leading term of the large  $\nu$  uniform asymptotic approximation (in R, the latter is available via function bessell.nuAsym() in package Bessel; Maechler 2012). If this is "sufficiently small" in absolute value (so that  $I_{\nu}(\kappa)$  will neither over- nor underflow), compute  $\log(I_{\nu}(\kappa))$  directly as the logarithm of  $I_{\nu}(\kappa)$ . Otherwise, if the approximation is "too large", but  $T_{\nu}(\kappa) - \kappa$  is sufficiently small in absolute value so that the exponentially scaled  $e^{-\kappa}I_{\nu}(\kappa)$  can be computed without over- or underflow, compute  $\log(I_{\nu}(\kappa))$  as  $\kappa + \log(e^{-\kappa}I_{\nu}(\kappa))$ . Otherwise, use the quick approximation, or the large  $\nu$  uniform asymptotic approximation with additional terms (package Bessel allows up to 4 additional terms). This strategy can readily be translated into a strategy for computing  $\log(H_{\nu})$ .

### 6. Application: useR! 2008 abstracts

In 2008 the "useR! 2008", the 3rd international R user conference, took place in Dortmund, Germany. In total 177 abstracts were submitted and accepted for presentation at the conference. The abstracts with additional information such as title, author, session, and keywords are available in the R data package corpus.useR.2008.abstracts available from the repository at <a href="http://datacube.wu.ac.at/">http://datacube.wu.ac.at/</a>.

The following code checks if the package is installed and if necessary installs it. Furthermore, the data contained in the package is loaded.

Using the **tm** package (Feinerer, Hornik, and Meyer 2008; Feinerer 2012) the data can be pre-processed by (1) generating a corpus from the vector of abstracts and (2) building a document-term matrix from the corpus. For constructing the document-term matrix each abstract needs to be tokenized (i.e., split into words, e.g., by using whitespace characters as separators) and transformed to lower case. Punctuation as well as numbers can be removed and the words can be stemmed (i.e., inflected words are reduced to their base form). In addition a minimum length can be imposed on the words as well as a minimum and maximum frequency inside an abstract required.

The vector of abstracts is transformed to an object which has an extended class of "Source" using VectorSource(). This object is used as input for Corpus() to generate the corpus. The map from the corpus to the document-term matrix is performed using DocumentTermMatrix(). The control argument of DocumentTermMatrix() specifies which pre-processing steps are applied to determine the frequency vectors of term occurrences in each abstract. We use the titles and the abstracts together to construct the document-term matrix.

```
R> library("tm")
R> abstracts_titles <-</pre>
```

Method "MC" was used for tokenizing. This method aims at producing the same results as MC (http://www.cs.utexas.edu/users/dml/software/mc/), which is a toolkit for creating vector models from text documents (Dhillon and Modha 2001). In addition the words are stemmed, a set of stopwords are removed and all words are kept which have a length of at least 3.

The resulting document-term matrix has 177 rows and 3853 columns. The ten most frequent terms (occurring in different abstracts) are the following.

```
R> library("slam")
R> ColSums <- col_sums(useR_2008_abstracts_DTM > 0)
R> sort(ColSums, decreasing = TRUE)[1:10]
     use
           packag
                       data
                                         model
                                                    base
                                                          develop
                                  can
     150
               127
                        121
                                  112
                                           103
                                                      91
                                                               91
 analysi
           method function
      88
               85
                         82
```

To reduce the dimension of the problem and omit terms which occur too frequently or too infrequently in the corpus to be of use when clustering the abstracts, we omit all terms which occur in less than 5 abstracts and more than 90 abstracts.

R> useR\_2008\_abstracts\_DTM <- weightTfIdf(useR\_2008\_abstracts\_DTM)

In the following different mixtures of vMF distributions are fitted to the training data when using 10-fold cross-validation and are compared based on the predictive log-likelihoods on the hold-out data to select a suitable model. The numbers of components are varied and the mixtures are fitted with concentration parameters constrained to be the same over components as well as where the concentration parameters are allowed to freely vary over components. For each training data set the EM algorithm is repeated 20 times with different random initializations.

```
R> set.seed(2008)
R> library("movMF")
R > Ks <- c(1:5, 10, 20)
R> splits <- sample(rep(1:10, length.out = nrow(useR_2008_abstracts_DTM)))
R> useR_2008_movMF <-
     lapply(Ks, function(k)
            sapply(1:10, function(s) {
              m <- movMF(useR_2008_abstracts_DTM[splits != s,],</pre>
                          k = k, nruns = 20)
              logLik(m, useR_2008_abstracts_DTM[splits == s,])
            }))
R> useR_2008_movMF_common <-
     lapply(Ks, function(k)
            sapply(1:10, function(s) {
              m <- movMF(useR_2008_abstracts_DTM[splits != s,],</pre>
                          k = k, nruns = 20,
                          kappa = list(common = TRUE))
              logLik(m, useR_2008_abstracts_DTM[splits == s,])
            }))
```

In Figure 1 the fitted models are compared using the cross-validated predictive log-likelihoods. The results for the models with free concentration parameters are on the left, for the models with common concentration parameters on the right. The predictive log-likelihoods on the hold-out data indicates that the best solutions have between 1 and 5 components and that the models with more components have rather bad predictive log-likelihoods.

Following the conclusions of the comparison of the predictive log-likelihoods values we further investigate the model where the concentration parameters are constrained to be equal over components and the number of components is equal to 2.

In the following we look at the 10 most important words of each fitted component:

```
R> apply(coef(best_model)$theta, 1, function(x)
+ colnames(coef(best_model)$theta)[order(x, decreasing = TRUE)[1:10]])

1     2
[1,] "user" "estim"
```

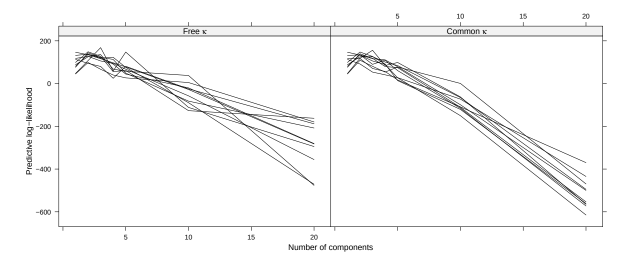


Figure 1: Predictive log-likelihoods for different and common concentration parameters  $\kappa$  for the fitted mixtures of vMF distributions to the "useR! 2008" abstracts.

```
[2,] "interfac" "variabl"
[3,] "project"
                  "method"
[4,] "gui"
                  "regress"
[5,] "analysi"
                  "function"
[6,] "statist"
                  "bayesian"
[7,] "system"
                  "robust"
[8,] "softwar"
                  "paramet"
[9,] "graphic"
                  "test"
[10,] "report"
                  "spatial"
```

Clearly one component deals with issues related to infrastructure or implementation, while the other component is focusing more on statistical modeling.

The clustering obtained from the a-posteriori probabilities is analyzed by comparing the cluster membership with the keywords assigned to an abstract. Because each abstract might have several keywords assigned, the abstracts and their cluster assignments are accordingly repeated.

In the following only keywords are shown which had more than 8 abstracts assigned to them.

```
R> tab <- tab[rowSums(tab) > 8,]
R> tab
```

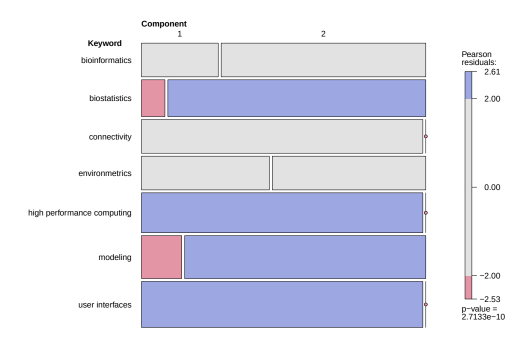


Figure 2: Cross-tabulation of the keywords in each session and the cluster assignment for keywords which were assigned to more than 8 abstracts.

(	Component		
Keyword	1	2	
bioinformatics	3	8	
biostatistics	1	11	
connectivity	11	0	
environmetrics	5	6	
high performance computing	13	0	
modeling	2	12	
user interfaces	15	0	

The table is also visualized in Figure 2. Abstracts where the keywords assigned relate to infrastructure or implementational issues such as connectivity, high performance computing and user interfaces are associated with one component, whereas abstracts which are related to statistical modeling issues such as bioinformatics, biostatistics and modeling are more likely to be assigned to the other component.

## 7. Summary

An R package for fitting finite mixtures of vMF distributions is presented. A separate package was developed in addition to general purpose mixture modeling packages in R such as for

example **flexmix** (Leisch 2004; Grün and Leisch 2008). This allows to take for example the special data structure into account. If this model class is applied in the context of text mining the input data will in general not only be high-dimensional, but also sparse. Using an implementation which allows to retain the sparse representation of the data therefore is crucial.

A possible extension for package **movMF** would be to allow for more flexible distributions in the components which also have as support the unit sphere. The vMF distribution is the analogue of the isotropic multivariate normal distribution, i.e., where the variance-covariance matrix is a multiple of the identity matrix. In  $\mathbb{R}^3$  the generalization of the vMF distribution which is the analogue to the general multivariate normal distribution on the two-dimensional unit sphere is the Fisher-Bingham (or Kent) distribution (Kent 1982). Finite mixtures of Fisher-Bingham distributions were used in Peel, Whiten, and McLachlan (2001) to identify joint sets present in rock mass. They also allowed for an additional component which followed the uniform distribution on the unit sphere. However, no generalization for higher dimensions are available and Dortet-Bernadet and Wicker (2008) propose to use inverse stereographic projections of multivariate normal distributions to cluster gene expression profiles.

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### Affiliation:

Kurt Hornik Institute for Statistics and Mathematics WU Wirtschaftsuniversität Wien Augasse 2–6 1090 Wien, Austria

E-mail: Kurt.Hornik@R-project.org

URL: http://statmath.wu.ac.at/~hornik/

Bettina Grün

Institut für Angewandte Statistik / IFAS

Johannes Kepler Universität Linz

Altenbergerstraße 69 4040 Linz, Austria

E-mail: Bettina.Gruen@jku.at URL: http://ifas.jku.at/gruen/