# fICA: FastICA Algorithms and Their Improved Variants

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**Abstract** In independent component analysis (ICA) one searches for mutually independent nongaussian latent variables when the components of the multivariate data are assumed to be linear combinations of them. Arguably, the most popular method to perform ICA is FastICA. There are two classical versions, the deflation-based FastICA where the components are found one by one, and the symmetric FastICA where the components are found simultaneously. These methods have been implemented previously in two R packages, **fastICA** and **ica**. We present the R package **fICA** and compare it to the other packages. The additional features of the package include for example optimization of the extraction order in the deflation-based version, possibility to use any nonlinearity function, and improvement to convergence of the deflation-based algorithm. The usage of the package is demonstrated by applying it to a real ECG data of a pregnant woman.

### Introduction

The goal of independent component analysis is to transform a multivariate dataset so that the resulting components are as independent as possible. The idea is to separate latent components from the dataset which is assumed to consist of linear mixtures of them.

The basic independent component (IC) model is thus written as

$$x_i = Az_i, i = 1, \dots, n, \tag{1}$$

where  $X = (x_1, ..., x_n)$  is a centered  $p \times n$  data matrix, A is a  $p \times p$  mixing matrix and  $z_1, ..., z_n$  are realizations of a zero mean random p-vector  $z = (z_1, ..., z_p)^T$  with mutually independent components, and at most one Gaussian component. The aim of ICA is then to estimate the unmixing matrix  $W = A^{-1}$  and/or Z from the data matrix X alone (Hyvärinen et al., 2001; Cichocki and Amari, 2002; Comon and Jutten, 2010; Nordhausen and Oja, 2018). Indeed, ensuring independence and nongaussianity of the components of z is sufficient for their consistent estimation, but only up to signs and scales (Comon, 1994). The scales of the independent components are often fixed by the assumption  $Cov(z) = I_p$ , and we may proceed from Model 1 to

$$x_{st} = U^T z, (2)$$

where  $x_{st} = \text{Cov}(x)^{-1/2}(x - \mathbb{E}(x))$  and U is an orthogonal  $p \times p$  matrix. This means the problem can be reduced to that of finding an orthogonal matrix and the final unmixing matrix estimate is then  $W = U\text{Cov}(x)^{-1/2}$ .

ICA has become a widely used tool in various fields, including brain imaging, image and audio signal processing and financial time series analysis. Despite the vast amount of literature on ICA methodology, the classical FastICA (Hyvärinen and Oja, 1997; Hyvärinen, 1999) methods are still the most popular tools to estimate the independent components with. FastICA functions are available for several programming languages such as Matlab: package FastICA (Gävert et al., 2005); R: packages fastICA (Marchini et al., 2013) and ica (Helwig, 2015); C++: part of package IT++ Cayre and Furon (2004); Python: part of packages MDP (Zito et al., 2008) and Scikit-learn (et al. Pedregosa, 2011).

Here we introduce the R package fICA (Miettinen et al., 2017b) which, in addition to the classical FastICA methods, includes functions for three recently proposed improved versions of FastICA, which are presented in the following section. Then we discuss how the fICA package differs from the packages fastICA and ica, and finally, we give some examples on the usage of the fICA package.

#### FastICA estimators in the fICA package

The **fICA** package includes implementations of the classical FastICA estimators as well as three improved variants, which we will now describe in detail. For other variants of FastICA, see for example Koldovský and Tichavský (2015).

All the methods have in common that they maximize the nongaussianity of the components of  $Ux_{st}$ , when the nongaussianity of a univariate random variable x is measured by  $|\mathbb{E}[G(x)]|$  with some twice continuously differentiable and nonquadratic function G which satisfies  $\mathbb{E}[G(y)] = 0$  for the standard Gaussian random variable y. The established and most popular options for G are

pow3: 
$$G(x) = (x^4 - 3)/4$$
,  
tanh:  $G(x) = log(cosh(x)) - c_t$ ,  
gaus:  $G(x) = -exp(-x^2/2) - c_g$ ,

where  $c_t = \mathbb{E}[\log(\cosh(y))] \approx 0.375$  and  $c_g = \mathbb{E}[-\exp(-y^2/2)] \approx -0.707$ . The names *pow3*, *tanh* and *gaus* originate from the first derivatives of the functions, the so called nonlinearities,  $g(x) = x^3$ ,  $g(x) = \tanh(x)$  and  $g(x) = x \exp(-x^2/2)$ . The choice *pow3* always yields consistent estimates, whereas *tanh* and *gaus* do not. However, the examples (Miettinen et al., 2017c; Wei, 2014) of such distributions of the independent components where *tanh* and *gaus* fail are quite artificial, and we agree with Hyvärinen (1999) stating that *tanh* and *gaus* work with most of the reasonable distributions. For further discussions concerning possible nonlinearities and their properties see for example Dermoune and Wei (2013) and Virta and Nordhausen (2017).

#### Deflation-based FastICA and its variants

The first FastICA paper (Hyvärinen and Oja, 1997) used *pow3* to find the independent components one after the other. In this *deflation-based FastICA*, the *k*th row of  $U = (u_1, ..., u_p)^T$  maximizes

$$|\mathbb{E}[G(\boldsymbol{u}_k^T\boldsymbol{x}_{st})]|$$

under the constraints  $u_k^T u_k = 1$  and  $u_j^T u_k = 0$ , j = 1, ..., k-1. A modified Newton-Raphson algorithm iterates the following steps until convergence.

$$u_k \leftarrow \mathbb{E}[g(u_k^T x_{st}) x_{st}] - \mathbb{E}[g'(u_k^T x_{st})] u_k$$

$$u_k \leftarrow \left(I_p - \sum_{l=1}^{k-1} u_l u_l^T\right) u_k$$

$$u_k \leftarrow ||u_k||^{-1} u_k.$$

The last two steps perform the Gram-Schmidt orthonormalization.

In addition to the global maximum, the objective function has several local maxima, and the FastICA algorithm may converge to the local maxima as well. The order in which the rows of U are found depends on their initial values. The extraction order affects the estimation performance, and to our knowledge, **fICA** is the only R package which provides a function to estimate the rows of U in such order that the objective function is maximized globally at each stage. This extraction order is usually better than a random order, even though not always optimal.

The statistical properties, e.g. asymptotic normality, of the deflation-based FastICA estimator were studied rigorously in Ollila (2009, 2010); Nordhausen et al. (2011). The derivation of the asymptotic variances of the unmixing matrix estimate lead to *reloaded FastICA* Nordhausen et al. (2011) which first computes an initial estimate  $\hat{Z}$  of the sources applying some simple IC method such as FOBI (Cardoso, 1989) or k-JADE (Miettinen et al., 2013). For all components of  $\hat{Z}$  one then computes certain quantities from which the optimal extraction order can be deduced.

The *adaptive deflation-based FastICA* (Miettinen et al., 2014) differs from the other estimators here, as instead of one nonlinearity it uses multiple nonlinearities by choosing for each component, out of a possible candidate set of nonlinearites, the best one. Again, an initial estimate of the sources is computed and then the same quantities as in reloaded FastICA are obtained, but now for for each candidate in the set of nonlinearities. Besides the optimal extraction order, also the optimal nonlinearity for each independent component separately is chosen. The default set of nonlinearities for the function adapt\_fICA in the R package fICA includes *pow3*, *tanh*, *gaus* and the following eleven:

$$\begin{array}{lll} g(x) = x^3 & g(x) = (x - 0.2)^2_+ + (x + 0.2)^2_- \\ g(x) = tanh(x) & g(x) = (x - 0.4)^2_+ + (x + 0.4)^2_- \\ g(x) = xexp(-x^2/2) & g(x) = (x - 0.6)^2_+ + (x + 0.6)^2_- \\ g(x) = (x + 0.6)^2_- & g(x) = (x - 0.8)^2_+ + (x + 0.8)^2_- \\ g(x) = (x - 0.6)^2_+ & g(x) = (x - 1.2)^2_+ + (x + 1.2)^2_- \\ g(x) = (x)^2_+ + (x)^2_- & g(x) = (x - 1.4)^2_+ + (x + 1.4)^2_- \\ g(x) = (x - 1)^2_+ + (x + 1)^2_- & g(x) = (x - 1.6)^2_+ + (x + 1.6)^2_- \end{array}$$

where  $(x)_+ = x$  if x > 0 and 0 otherwise, and  $(x)_- = x$  if x < 0 and 0 otherwise.

#### Symmetric and squared symmetric FastICA

The symmetric FastICA (Hyvärinen, 1999) estimator maximizes

$$\sum_{i=1}^{p} | \mathbb{E}[G(\boldsymbol{u}_{j}^{T} \boldsymbol{x}_{st}] |$$

under the orthogonality constraint  $UU^T = I_p$ . Now the rows of  $U = (\mathbf{u}_1, \dots, \mathbf{u}_p)^T$  are estimated in parallel, and the steps of the iterative algorithm are

$$u_k \leftarrow \mathbb{E}[g(u_k^T x_{st}) x_{st}] - \mathbb{E}[g'(u_k^T x_{st})] u_k, \quad k = 1, \dots, p$$
  
$$U \leftarrow (UU^T)^{-1/2} U.$$

The first update step is similar to that of the deflation-based version. In a way, the orthogonalization step can be seen as taking an average over the vectors of the first step, while in the Gram-Schmidt orthogonalization the errors in estimating the first rows of U will remain throughout the estimation. It is said that the error accumulates in the deflation-based approach. The symmetric FastICA is usually considered superior to the deflation-based FastICA, but there are also cases where the accumulation is preferable to the averaging. This occurs when some independent components are easier to find than the others. Statistical properties of symmetric FastICA are given in Miettinen et al. (2015); Wei (2015); Miettinen et al. (2017c).

The squared symmetric FastICA (Miettinen et al., 2017c) estimator maximizes

$$\sum_{j=1}^{p} (\mathbb{E}[G(\boldsymbol{u}_{j}^{T}\boldsymbol{x}_{st}])^{2}$$

under the orthogonality constraint  $UU^T = I_p$ . The first step of the algorithm

$$u_k \leftarrow \mathbb{E}[G(u_k^T x_{st})](\mathbb{E}[g(u_k^T x_{st}) x_{st}] - \mathbb{E}[g'(u_k^T x_{st})]u_k), \ k = 1, \dots, p,$$
  
$$U \leftarrow (UU^T)^{-1/2}U$$

is that of the classical symmetric version multiplied by  $\mathbb{E}[G(u_k^Tx_{st})]$ . Hence, the squared symmetric version puts more weight on the rows which correspond to more nongaussian components, which most often, but not always, is advantageous.

For both of the symmetric versions, the initial value of U is not important when the sample size is large, as the algorithms converge always to the global maxima. With small sample sizes the initial value plays a role.

# Comparison of the packages

The FastICA algorithms sometimes fail to converge when the sample size is small. However, this is not seen if one uses the R package **fastICA** or **ica**, since their functions do not give errors or warnings if the algorithm does not converge, but simply return the estimate after the maximum number of iterations is reached. In the **fastICA** package, the information about convergence can be obtained, but since it is not given in the default setup, many of the users will not notice it. Similarly, in the **ica** package the number of iterations is included in the output, and it could be used as an indicator for convergence, but problems might often still go unnoticed. Growing the number of iterations does usually not help because most often when the algorithm has not converged until 2000 iterations, the problem is that the algorithm gets stuck and starts to repeat a loop of values. The functions in the R package **fICA** give an error message if the algorithm does not converge. When computing the symmetric estimators, the user can choose the number of random orthogonal matrices which are generated for the initial values of the algorithms. If this number is at least two, the function returns the estimate which yields the highest value of the objective function, and gives a warning if none of the algorithm runs converge. To avoid the repeating loop in the deflation-based case, the function in the **fICA** package alternates the step size, that is, the update of the vector  $u_k$  is of the form

$$u_k \leftarrow w u_k + (1 - w) \left( \mathbb{E}[g(u_k^T x_{st}) x_{st}] - \mathbb{E}[g'(u_k^T x_{st})] u_k \right),$$

where w is zero for most iterations and takes small nonzero values with some selected iteration numbers with an irregular pattern.

To sum up the key differences of **fICA** to the **fastICA** and **ica** packages, we present Table 1.

Table 1: The properties of the three R packages which include the FastICA algorithm.

	fICA	fastICA	ica
Symmetric FastICA	YES	YES	YES
Squared symm. FastICA	YES	NO	NO
Deflation-based FastICA	YES	YES	YES
Optimized extraction order	YES	NO	NO
Adaptive defl. FastICA	YES	NO	NO
Error if no convergence	YES	NO	NO
Number of nonlinearities	$\infty$	2	3

The number of nonlinearities is in the **fICA** basically unlimited as besides the already implemented suggestions the user can provide own functions by specifying g and its derivative.

## **Examples**

To illustrate the use of the fICA package, we give the following two examples.

#### Electrocardiography data

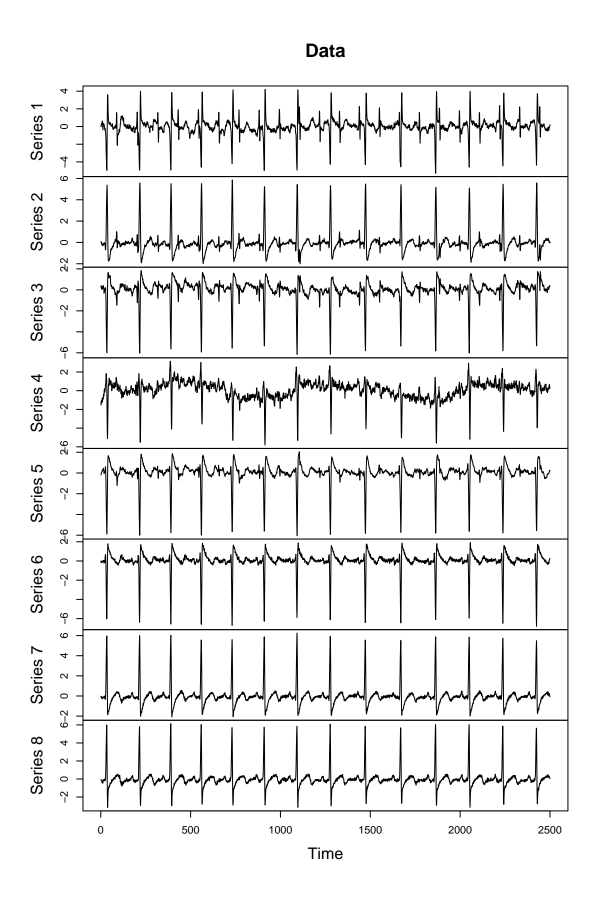
First, we analyze an electrocardiography recording (ECG) dataset 'foetal\_ecg.dat' ( de Lathauwer et al., 2000), which can be downloaded from the supplementary files of Miettinen et al. (2017d). ECG measures the electrical potential, which is mainly generated by the heart beats, on the body surface. This recording is from a pregnant woman, and it consists of eight signals, the five first of them measured by sensors on the stomach and the three last measured on the chest. The lengths of the signals are 2500 time points.

The signals are mixtures of fetus' and mother's heart beats and possibly some artifacts, and the goal is to separate these sources using independent component analysis.

```
> library(fICA)
> library(JADE)
> options(digits = 4)
>
> dataset <- matrix(scan("https://www.jstatsoft.org/index.php/jss/article/
+ downloadSuppFile/v076i02/foetal_ecg.dat"), 2500, 9, byrow = TRUE)
> X <- dataset[ , 2:9]
> plot.ts(X, nc = 1, main = "Data")
```

After downloading the data and plotting it in Figure 1, we see that the main feature in each signal are 14 peaks, which are presumably produced by mother's heart beats. Next we show how to apply the function adapt\_fICA with a modified set of nonlinearities, which includes pow3, tanh and gaus which can be found from the package, and another classical nonlinearity called skew given by  $g(x) = x^2$ . As the nonlinearity skew is not implemented the user can add it simply by providing functions for g and its derivative.

```
> g \leftarrow function(x) \{ x^2 \}
> dg \leftarrow function(x) \{ 2*x \}
> gf_new <- c(gf[1:3], g)
> dgf_new <- c(dgf[1:3], dg)
> gnames_new <- c(gnames[1:3], "skew")</pre>
> res <- adapt_fICA(X, gs = gf_new, dgs = dgf_new, name = gnames_new, kj = 1)</pre>
>
> res
W :
                                    [,4]
        [,1]
                 [,2]
                          [,3]
                                             [,5]
                                                       [,6]
                                                                  [,7]
[1,] -0.04135  0.059521  0.004006  0.001754 -0.010123  0.0076023  0.0001551
[2,] 0.01731 -0.023294 0.002298 0.002663 0.008367 0.0022727 -0.0075911
[3,] 0.09352 0.071699 -0.106091 -0.004342 0.175369 -0.0030657 -0.0063677
[5,] 0.15805 0.039095 0.233377 -0.041272 -0.169486 0.0104065 -0.0116267
```



**Figure 1:** The ECG signals of a pregnant woman.

```
[6,] -0.06605 0.114817 0.201392 -0.021387 0.234210 -0.0106505 0.0254791
[7,] 0.07817 -0.001272 0.146718 0.220175 -0.180189 -0.0008764 0.0035999
[8,] -0.25492  0.313232  0.049493  0.089799  0.028816  0.0232881 -0.0535985
         [,8]
[1,] -0.009234
[2,] 0.009301
[3,] 0.013034
[4,] 0.022396
[5,] 0.028662
[6,] -0.010261
[7,] -0.009922
[8,] 0.034743
gs:
[1] "pow3" "tanh" "gaus" "skew"
used gs:
[1] "gaus" "gaus" "tanh" "tanh" "tanh" "tanh"
alphas :
    comp 1 comp 2 comp 3 comp 4 comp 5 comp 6 comp 7 comp 8
pow3 0.7445 0.6119 1.4060 1.994 5.014 9.628 23.28 50087.8
tanh 0.2433 0.2569 0.7769 1.374 3.189 8.400 21.53 343.3
gaus 0.2058 0.2336 0.7336 1.458 3.351 10.829 22.87
                                                      215.1
skew 0.5457 0.4126 11.8980 7.091 12.867 9.118 25.00
                                                     108.7
init_est :
Γ17 "1-JADE"
```

In the output we have the  $8 \times 8$  unmixing matrix, the names of the nonlinearities which were available, the nonlinearities which were used, the criterion values for the selection of the extraction order and the nonlinearities, and the name of the initial estimator.

Note that the used\_gs does not give a nonlinearity for the last component, because the last component is fixed by the seven previous components due to the orthogonality constraint.

```
> plot.ts(bss.components(res), nc=1, main="Estimated components")
```

Figure 2 plots the estimated independent components. Clearly, the first two components are related to mother's heart beat and the third one is related to fetus' heart beat. The third row of the unmixing matrix estimate shows which data signals contribute to the estimate of the fetal heart beat. Not surprisingly, the nonzero coefficients correspond to sensors 1, 2, 3 and 5, which are located on the stomach area.

#### **Simulations**

In this simulation example, we also take a look at the related R package BSSasymp (Miettinen et al., 2017a), which computes asymptotic covariance matrices of several mixing and unmixing matrix estimates, including the FastICA variants discussed in this paper and implemented in fICA. The p=3 independent components in the simulation setup are generated from a  $t_9$ -distribution, an exponential distribution with rate 1, and the normal distribution. Each density is standardized so that the expected value is 0 and the variance is 1. In each of the 1000 repetitions, the sample size is 5000, the mixing matrix is the same randomly generated  $3 \times 3$  matrix, and we collect the unmixing matrix estimates given by (1) the reloaded deflation-based FastICA using tanh from fICA, (2) deflation-based FastICA using tanh from from faca. The goal of the simulation is to show that the extraction order of the components indeed matters.

```
> library(fICA)
> library(fastICA)
> library(ica)
> library(BSSasymp)
> library(JADE)
>
> options(digits = 4)
> set.seed(1145)
```

# **Estimated components**

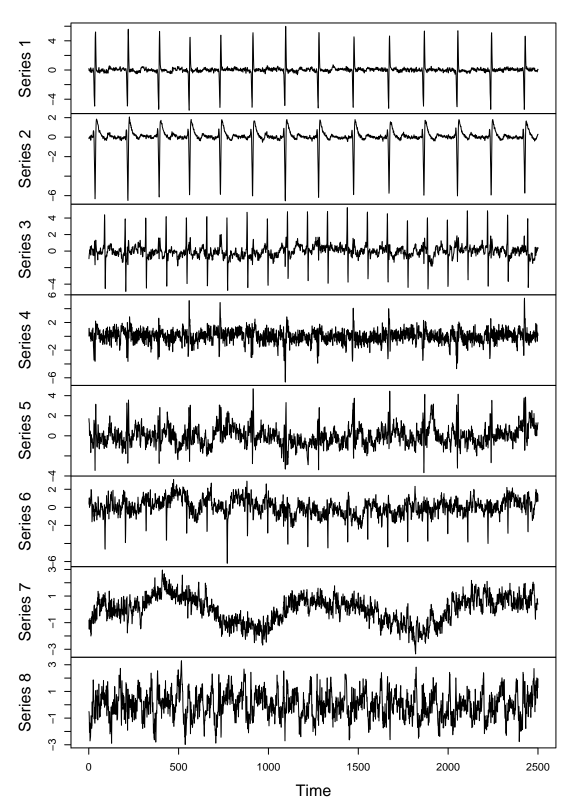


Figure 2: The estimated independent components from the ECG data.

```
> rort <- function(p){qr.Q(qr(matrix(rnorm(p * p), p)))}</pre>
>
> n <- 5000
> p <- 3
> A <- matrix(rnorm(p^2), p, p)</pre>
> Ws1 <- Ws2 <- Ws3 <- vector("list", 1000)
> for(i in 1:1000){
    Z \leftarrow cbind(rt(n, 9) / sqrt(9/7), rexp(n, 1) - 1, rnorm(n))
    X \leftarrow tcrossprod(Z,A)
+
+
    init = rort(p)
    Ws1[[i]] <- reloaded_fICA(X, g = "tanh")$W</pre>
    res <- fastICA(X, n.comp = 3, alg.typ = "deflation" )
    Ws2[[i]] <- t(res$W) %*% t(res$K)
    Ws3[[i]] <- icafast(X, nc = 3, alg = "def", Rmat = init)$W
+ }
```

To obtain the theoretical asymptotic variances, the density functions of the independent components, and their supports are required. Function ASCOV\_FastICAdefl computes the values for deflation-based variants of FastICA. However, in the case that the initial matrix is random, as it is in fastICA and ica, there are no asymptotic results, and therefore next we only consider the reloaded FastICA, for which the asymptotic variances are derived using ASCOV\_FastICAdefl with method = "adapt" and only one nonlinearity. The output component \$COV\_W gives the asymptotic covariance matrix of the unmixing matrix estimate, and the variances are then on the diagonal column after column. The variances can be estimated using the function ASCOV\_FastICAdefl\_est and the data matrix. Here we estimate the variances from the last dataset generated above. In this case, the estimated variances are slightly higher than the theoretical values. Finally, we compute the variances from the simulations, and notice that they are quite close to the theoretical and the estimated variances.

```
> f1 <- function(x) \{ dt(x*sqrt(9/7),9)*sqrt(9/7) \}
 > f2 <- function(x){ dexp(x+1,1) }
 > f3 <- function(x){ dnorm(x) }</pre>
 > supp <- matrix(c(-Inf, -1, -Inf, Inf, Inf, Inf), p, 2)
> COV <- ASCOV_FastICAdefl(c(f1,f2,f3), gf[2], dgf[2], Gf[2],</pre>
 + method = "adapt", name = c("tanh"), supp = supp, A = A)$COV_W
 > matrix(diag(COV), p, p)
      [,1] [,2] [,3]
[1,] 6.974 1.486 4.157
[2,] 27.813 5.163 10.895
[3,] 4.703 1.533 9.879
> COV_est <- ASCOV_FastICAdefl_est(X, gf[2], dgf[2], Gf[2],</pre>
+ method="adapt", name=c("tanh"))$COV_W
> matrix(diag(COV_est), p, p)*n
       [,1] [,2] [,3]
[1,] 7.219 1.369 4.356
[2,] 39.321 6.399 12.983
[3,] 5.714 1.552 16.177
> apply(simplify2array(Ws1), c(1,2), var)*n
       [,1] [,2] [,3]
[1,] 7.128 1.582 4.132
[2,] 29.009 5.334 11.498
[3,] 4.939 1.528 10.639
```

The minimum distance index (Ilmonen et al., 2010) can be used to quantify the separation performance of an unmixing matrix  $\hat{W}$  in a simulation study where the mixing matrix is known. It is defined as

$$\hat{D} = \frac{1}{\sqrt{p-1}} \inf_{C \in \mathcal{C}} \|C\widehat{W}A - I_p\|,$$

where  $\|\cdot\|$  is the Frobenius norm and

```
C = \{C : \text{ each row and column of } C \text{ has exactly one non-zero element}\}.
```

The minimum distance index takes values between 0 and 1, where zero means perfect separation. The function ASCOV\_FastICAdef1 gives the limiting expected value of  $\hat{D}^2n(p-1)$ , which is independent of the mixing matrix A. Below, this expected value is computed for the optimal extraction order (first exponential, then  $t_9$  distribution and last the normal component), and for the other reasonable order, where the  $t_9$ -distributed component is found first, the exponentially distributed second, and the Gaussian last. If the Gaussian component is found first or second, the limiting expected value goes to infinity. We also compute the averages of  $\hat{D}^2n(p-1)$  over the 1000 repetitions for the three estimates. The average of the reloaded FastICA is close to the expected value of the optimal order, while the other two estimates with a random initial value, have larger averages as the usually find the components in the second best order.

```
> EMD_opt <- ASCOV_FastICAdefl(c(f1,f2,f3), gf[2], dgf[2], Gf[2],</pre>
+ method = "adapt", name = c("tanh"), supp = supp, A = A)$EMD
> EMD_opt
[1] 44.74
> EMD2 <- ASCOV_FastICAdefl(c(f1,f2,f3), gf[2], dgf[2], Gf[2],</pre>
+ method = "regular", name = c("tanh"), supp = supp, A = A)$EMD
> EMD2
[1] 67.66
> MD1 <- unlist(lapply(Ws1, MD, A=A))
> MD2 <- unlist(lapply(Ws2, MD, A=A))</pre>
> MD3 <- unlist(lapply(Ws3, MD, A=A))</pre>
> mean(MD1^2)*n*(p-1) # fICA
Γ17 46.74
> mean(MD2^2)*n*(p-1) # fastICA
[1] 67.87
> mean(MD3^2)*n*(p-1) # ica
[1] 69.34
```

Hence this simulation demonstrations shows clearly the effect of the extraction order in deflationbased fastICA and that only **fICA** has tools to find the optimal order.

#### Summary

FastICA is the most widely used method to carry out independent component analysis. The R package **fICA** contributes to CRAN in two ways. First, it introduces the reloaded and adaptive deflation-based FastICA and the squared symmetric FastICA, and second, it improves on the existing R functions for classical FastICA algorithms by allowing user-specified nonlinearities and enhancing the convergence of the deflation-based FastICA. We gave a short review of the FastICA estimators, and examples on how to use them with the **fICA** package.

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