extTADA for developmental disorder (DD)

This example describes steps to obtain results for DD de novo data.

Some main steps:

- 1. Estimate genetic parameters using MCCMC.
- 2. Use these parameters to calculate FDRs for each gene.

Load source files of extTADA

```
In [1]: fileR <- dir("../script", ".R$")</pre>
        for (ii in fileR)
            source(paste0("../script/", ii))
        Warning message:
        : package 'rstan' was built under R version 3.2.5Loading required package: ggplot2
        Warning message:
        : package 'ggplot2' was built under R version 3.2.4Loading required package: StanHeaders
        Warning message:
        : package 'StanHeaders' was built under R version 3.2.5rstan (Version 2.12.1, packaged: 2016-09-11
        13:07:50 UTC, GitRev: 85f7a56811da)
        For execution on a local, multicore CPU with excess RAM we recommend calling
        rstan_options(auto_write = TRUE)
        options(mc.cores = parallel::detectCores())
        locfit 1.5-9.1 2013-03-22
In [2]: data <- read.table("../data/data_mut_DD.csv", header = TRUE, as.is = TRUE)</pre>
        head(data)
```

Out[2]:

		Gene	mut_lof	mut_missense	mut_damaging	dn_damaging_DD	dn_lof_DD	dn_missense_DD	dn_silent_DD
	1	A1BG	9.996657e- 07	2.300224e-05	4.54062e-07	0	0	0	0
	2	A1BG- AS1	1.420491e- 07	5.566513e-08	1.04146e-10	0	0	0	0
;	3	A1CF	2.144318e- 06	1.68827e-05	3.35199e-06	0	0	0	0
	4	A2M	3.981797e- 06	4.039597e-05	7.82154e-06	0	0	0	0
,	5	A2M-AS1	9.14698e-08	5.566513e-08	1.04146e-10	0	0	0	0
	6	A2ML1	3.779929e- 06	4.007178e-05	2.85364e-06	0	0	1	0

```
In []:

In [3]: allDNData <- data[, paste0("dn_", c("damaging", "lof"), "_DD")]
      allMutData <- data[,paste0("mut_", c("damaging", "lof"))]
      head(data.frame(allMutData, allDNData))</pre>
```

Out[3]:

	mut_damaging	mut_lof	dn_damaging_DD	dn_lof_DD
1	4.54062e-07	9.996657e-07	0	0
2	1.04146e-10	1.420491e-07	0	0
3	3.35199e-06	2.144318e-06	0	0
4	7.82154e-06	3.981797e-06	0	0
5	1.04146e-10	9.14698e-08	0	0
6	2.85364e-06	3.779929e-06	0	0

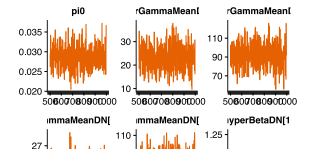
Use the function extTADA to sample values of parameters

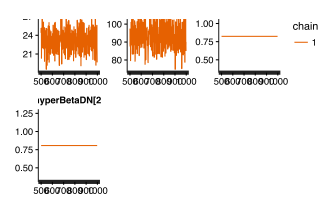
```
Ndn = rep(4293, 2),
                  nIteration = 1000,
                  nIteration2 = 2000)
There are 19358 in this analysis
Sampling with nter = 1000 and nThin = 1
The model DNextTADA is used
clang: warning: optimization flag '-ffat-lto-objects' is not supported
clang: warning: argument unused during compilation: '-ffat-lto-objects'
SAMPLING FOR MODEL '493590f0b50b55273a713acc3c2b8395' NOW (CHAIN 1).
Chain 1, Iteration:
                     1 / 1000 [ 0%]
                                       (Warmup)
Chain 1, Iteration: 100 / 1000 [ 10%]
                                       (Warmup)
Chain 1, Iteration: 200 / 1000 [ 20%]
                                       (Warmup)
Chain 1, Iteration: 300 / 1000 [ 30%]
Chain 1, Iteration: 400 / 1000 [ 40%]
                                       (Warmup)
Chain 1, Iteration: 500 / 1000 [ 50%]
                                       (Warmup)
Chain 1, Iteration: 501 / 1000 [ 50%]
                                       (Sampling)
Chain 1, Iteration: 600 / 1000 [ 60%]
                                       (Sampling)
Chain 1, Iteration: 700 / 1000 [ 70%]
                                       (Sampling)
Chain 1, Iteration: 800 / 1000 [ 80%]
                                       (Sampling)
Chain 1, Iteration: 900 / 1000 [ 90%]
                                       (Sampling)
Chain 1, Iteration: 1000 / 1000 [100%] (Sampling)
Elapsed Time: 43.3239 seconds (Warm-up)
               32.5705 seconds (Sampling)
               75.8945 seconds (Total)
Sampling with nter = 2000 and nThin = 2
The model DNextTADA is used
clang: warning: optimization flag '-ffat-lto-objects' is not supported
clang: warning: argument unused during compilation: '-ffat-lto-objects'
SAMPLING FOR MODEL '493590f0b50b55273a713acc3c2b8395' NOW (CHAIN 1).
Chain 1, Iteration:
                       1 / 2000 [ 0%]
                                        (Warmup)
Chain 1, Iteration: 200 / 2000 [ 10%]
                                        (Warmup)
Chain 1, Iteration: 400 / 2000 [ 20%]
                                        (Warmup)
Chain 1, Iteration: 600 / 2000 [ 30%]
                                        (Warmup)
Chain 1, Iteration: 800 / 2000 [ 40%]
                                        (Warmup)
Chain 1, Iteration: 1000 / 2000 [ 50%]
                                        (Warmup)
Chain 1, Iteration: 1001 / 2000 [ 50%]
                                        (Sampling)
Chain 1, Iteration: 1200 / 2000 [ 60%]
                                        (Sampling)
Chain 1, Iteration: 1400 / 2000 [ 70%]
                                        (Sampling)
Chain 1, Iteration: 1600 / 2000 [ 80%]
                                        (Sampling)
Chain 1, Iteration: 1800 / 2000 [ 90%]
                                        (Sampling)
Chain 1, Iteration: 2000 / 2000 [100%] (Sampling)
Elapsed Time: 75.5381 seconds (Warm-up)
               68.6742 seconds (Sampling)
               144.212 seconds (Total)
```

dataDN = allDNData, mutRate = allMutData,

Take a quick look at the traces of parameters

```
In [5]: options(repr.plot.width=5, repr.plot.height=5)
    stan_trace(mcmcDD)
```





```
In [6]: mcmcDD
```

Out[6]: Inference for Stan model: 493590f0b50b55273a713acc3c2b8395.

1 chains, each with iter=2000; warmup=1000; thin=2;
post-warmup draws per chain=500, total post-warmup draws=500.

	mean	se mean	sd	2.5%	25%	50%	75%
pi0	0.03	0.00	0.00	0.02	0.03	0.03	0.03
hyperGammaMeanDN[1]	23.21	0.27	5.02	13.55	19.81	23.38	26.53
hyperGammaMeanDN[2]	92.04	0.65	12.34	69.32	83.19	92.65	100.08
<pre>gammaMeanDN[1]</pre>	23.12	0.08	1.60	20.03	22.06	23.05	24.09
gammaMeanDN[2]	93.57	0.37	6.61	81.51	88.92	93.21	97.91
hyperBetaDN[1]	0.82	0.00	0.00	0.82	0.82	0.82	0.82
hyperBetaDN[2]	0.81	0.00	0.00	0.81	0.81	0.81	0.81
lp	-6792.93	0.07	1.49	-6796.77	-6793.69	-6792.59	-6791.82
	97.5%	n_eff Rh	nat				
pi0	0.03	311 1	.00				
hyperGammaMeanDN[1]	32.59	338 1	.00				
hyperGammaMeanDN[2]	114.98	361 1.	.00				
<pre>gammaMeanDN[1]</pre>	26.13	409 1	.00				
gammaMeanDN[2]	105.68	317 1	.00				
hyperBetaDN[1]	0.82	1 1	.00				
hyperBetaDN[2]	0.81	1 1	.00				
lp	-6791.03	423 1	.01				

Samples were drawn using NUTS(diag_e) at Wed Dec 14 16:00:11 2016. For each parameter, n_eff is a crude measure of effective sample size, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat=1).

Use the function estimatePars of extTADA to obtain modes, and credible intervals (ID) of parameters

==== Only pi and hyper parameters are estimated in this step extTADA does not calculate HPDs for hyper betas, just their medians ===

In [8]: pars0

Out[8]:

	Mode	ICI	uCl
pi0	0.02870958	0.02327248	0.03431055
hyperGammaMeanDN[1]	23.60775	13.52292	32.87455
hyperGammaMeanDN[2]	93.46348	68.64038	114.97203
hyperBetaDN[1]	0.8232544	0.8232544	0.8232544
hyperBetaDN[2]	0.8066756	0.8066756	0.8066756

Use the function plotParHeatmap of extTADA to draw heatmaps of pairs of pars

```
In [9]: Options(repr.piot.wiatn=4, repr.piot.neignt=3)
          par(mfrow = c(1, 2))
          plotParHeatmap(pars = c("pi0", "hyperGammaMeanDN[1]"), mcmcResult = mcmcDD)
         plotParHeatmap(pars = c("pi0", "hyperGammaMeanDN[2]"), mcmcResult = mcmcDD)
         Warning message:
         In plot.xy(xy, type, ...): font width unknown for character 0x1Warning message:
         In plot.xy(xy, type, ...): font metrics unknown for character 0x1Warning message:
         In plot.xy(xy, type, ...): font width unknown for character 0x1Warning message: In plot.xy(xy, type, ...): font metrics unknown for character 0x1
          hyperGammaMeanDN[1]
                                     hyperGammaMeanDN[2]
                8
                                           8
                                           4
                9
                   0.00
                           0.03
                                              0.00
                                                      0.03
```

pi0

Use function calculateFDR of extTADA to obtain FDRs of genes

pi0

No parameters for case-control data; therefore, these categories are not calculated in this step.

In [14]: head(dataFDR)

Out[14]:

	geneName	dn_damaging_DD	dn_lof_DD	mut_damaging	mut_lof	BF	qvalue
681	ANKRD11	0	32	1.12154e-05	4.918288e-06	2.153447e+62	0
1001	ARID1B	0	30	9.81608e-06	4.623958e-06	2.411403e+58	0
10146	MLL	1	26	1.19316e-05	9.084711e-06	1.877296e+49	0
347	ADNP	1	19	2.26581e-06	2.100249e-06	4.275545e+38	0
4832	DYRK1A	4	14	4.53477e-06	2.145461e-06	2.406051e+32	0
9906	MED13L	5	13	1.5154e-05	4.661034e-06	5.968457e+29	0

```
In [15]: dim(dataFDR[dataFDR$qvalue < 0.1, ])
    dim(dataFDR[dataFDR$qvalue < 0.05, ])
Out[15]: 1.199</pre>
```

```
Out[15]: 1. 199
2. 7
Out[15]: 1. 162
2. 7
```