1	Manual for Euroformix v1:
2	(A) Installation and maning magazine
3	(A) <u>Installation and running program:</u>
4 5	1) Run R (>=3.0.1) in Windows, Linux or MAC (http://cran.r-project.org/).
6	2) Required packages to run GUI:
7	a. gWidgetstcltk (depends on digest,tcltk)
8	b. gWidgets
9 10	Other required packages: a. cubature
11	i. Required for multivariate integration (Integrated LR).
12	b. forensim
13	i. Required for qualitative Weight-of-Evidence.
14	4) Installation and run gammadnamix:
15 16	a. install.packages("gammadnamix", repos="http://R-Forge.R-project.org")b. library(gammadnamix)
17	c. euroformix()
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25	(D) CIII
26	(B) GUI
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28	Sections:
29	0- <u>Toolbar</u>
30	1- <u>Importing data</u>
31	2- Model specification
32	3- MLE fit: ('Continuous LR (Maximum Likelihood based)')
33	4- Deconvolution (Deconvolution based on the continuous model)
34	5- Database Search (Database search based on the continuous and
35	qualitative model)
36	6- Qual.LR (Qualitative model)
37	7- Generate data (Generation from the continuous model)
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39	0. <u>Toolbar</u>
40	- File
41	• Set directory: The user may select the working directory of the R-program.
42	 Open project: The user may open an earlier project which is saved in a file on the form
43	"projectname.Rdata".
44	 Save project: The user may save the existing project into a file with name
45	"project name".
46	Extension .Rdata is added automatically to project name.
47	 All data imported to the program and resulting calculations are stored into a
48	single project-file which may be open at any time in the program.
49	Saving a project makes:
50	Big reference databases are stored efficiently (the required space for the
51	database is drastically reduced).
52	• Time-consuming calculations are restored instantly (only required to be
53	calculated ones).
54	- Frequencies
55	 Set size of frequency database: User may specify number of samples 'N' used to create
56	the population frequencies.
57	 When new alleles from imported files are found, these are assigned as freq0.
58	• If N=0 (this is default), freq0 is equal minimum observed frequency.
59	• If N>0, freq0='5/(2N)'.
60	New alleles are updated to the population frequencies when:
61	 When a reference database is imported.
62	 When interpretations are done.
63	 Deconvolution, Weight-of-Evidence and 'Database search'
64	 Frequencies are normalized for each of these two cases.
65	• WARNING: Normalizing may be done twice if new alleles (not
66	seen in population frequency table or reference database) are
67	observed in the evidence/reference profile.
68	 Set number of wildcards in false positive match: The user may specify number of
69	wildcards in the random match probability statistics
70	- Optimization
71	 Set number of random startpoints: The user may set required number of independent
72	random startpoints in the optimizer to ensure that the global maximum is attained for the
73	Maximum Likelihood Estimator (MLE). Default is 3.
74	 Set variance of randomizer: The user may set the variance parameter used for the
75	random generation of startpoints used in optimizer. Default is 10.
76	- MCMC (Markov Chain Monte Carlo)
77	 Set number of samples: The user may set the number of samples drawn from the
78	posterior distribution of the parameters. Default is 10000.
79	o Set variance of randomizer : The user may set the variance parameter scalar used in the
80	'Markov Chain Monte Carlo (MCMC) random walk Metropolis'. See vignette for
81	details. Default is 10.
82	 Note that this value should be tweaked such that acceptance rate of sampler are
83	around 0.2 (to ensure global exploration in the parameter space).

- Integration
 - **Set relative error requirement:** The user may set the required estimated relative error used in the integration function adaptIntegrate {cubature}. See vignette for details. Default is 0.005.
 - o **Set maximum of mu-parameter**: The user may set upper limit of mu-parameter (amount of DNA). See vignette for details. Default is 20000.
 - **Set maximum of sigma-parameter**: The user may set upper limit of sigma-parameter (coefficient of variation). See vignette for details. Default is 1.
 - **Set maximum of stutter ratio-parameter**: The user may set upper limit of the stutter ratio parameter (xi). Default is 1.
- Deconvolution
 - **Set required summed probability**: The user may set required summed posterior genotype-probability which the deconvoluted list is ensured to contain. Default is 0.9999.
 - **Set max listsize:** The user may set maximum number of elements in the deconvoluted list. Default is 1000.
 - The greater max listsize, the more time-consuming (and memory consuming) the search-algorithm behind will be.
- Database search
 - o **Set maximum view-elements**: The user may set maximum number of individuals to show from the reference-database. Default is 10000.
 - The greater 'value', the more time-consuming will it become to show table on screen.
 - Note that the result table from the database search shows only the top 'value'ranked elements.
 - O Set drop-in probability for qualitative model: When searching database with continuous LR model, the qualitative LR model is also considered with a specific dropin probability parameter given here (default is 0.05).

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1. Importing data

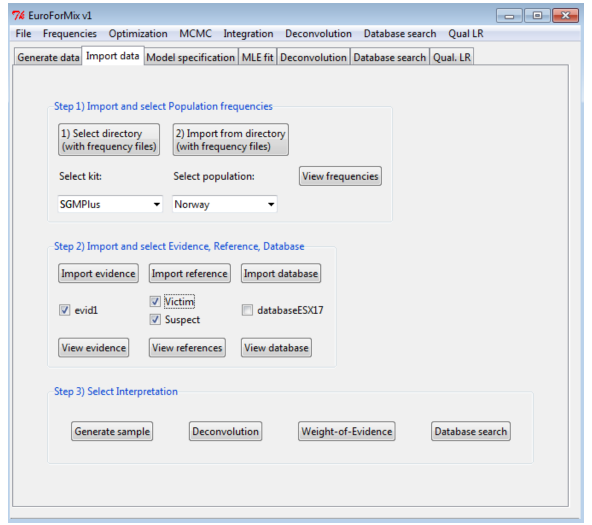


Figure 1: The figure shows the <u>Import data</u> GUI page where the user can import population frequencies, evidence stains, reference profiles and reference databases.

DATA IMPORT:

- **Common** for all files:
 - The extension (denotes file-type) of the file names does not matter. It may also have no extension at all.
 - All imported files must be either comma, semi-colon or tab-separated (',',',','\t').
 - o Required/optional headers (all are capital invariant):
 - "sample" is required header for sample(s) name(s).
 - The sample names are NOT capital invariant.
 - If more than one header name contains "sample", it will select the header name which in addition contains "name" in the same string.
 - "marker" is required header for marker name(s).

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- Marker names are capital invariant.
- If no header is found, the header containing "loc" will be used if found.
- "allele" is required header(s) for allele-information.
 - This may be a vector ("alleleX1",..., "allelleX10") of any length denoting allele(s) to a given marker for a given sample. Here X1,...,X10 can be anything.
- "height" optional header(s) for peak height-information.
 - This may be a vector ("heightX1",..., "heightX10") of any length denoting peak height to the corresponding allele(s) in "allele". Here X1,...,X10 can be anything.

o Note:

- The imported data will use upper-letter of marker-names found in the file.
- All imports are printed out in the terminal (see figure 2). From this, the user may check that the data are imported correctly.

[1] "Raw fil i	mport:"								
	Sample.Name	Marker	Allele.1	Allele.2	Allele.3	Alle	le.4	Allele.5	Allele.6	Height.1
1	evid1	AMEL	X	Y	NA		NA	NA	NA	2136
2	evid1	D3S1358	14	15	16.0	1	NA	NA	NA	178
3	evid1	TH01	6	7	9.3		NA	NA	NA	419
4	evid1	D21S11	27	29	NA		NA	NA	NA	1128
5	evid1	D18S51	15	17	NA	L	NA	NA	NA	467
6	evid1	D2S1338	17	19	20.0		23	NA	NA	290
7	evid1	D16S539	9	10	11.0	1	12	NA	NA	217
8	evid1	vWA	14	15	17.0		NA	NA	NA	1250
9	evid1	D8S1179	10	13	14.0	1	15	NA	NA	206
10	evid1	FGA	21	22	NA	L	NA	NA	NA	664
11	evid1	D19S433	13	14	15.2		NA	NA	NA	1157
	Height.2 He	ight.3 H	eight.4 H	eight.5 H	eight.6	ADO	UD1	X		
1	1015	NA	NA	NA	NA f	alse	NA :	NA		
2	2405	1982	NA	NA	NA f	alse	NA :	AN		
3	282	1871	NA	NA	NA f	alse	NA :	NΑ		
4	1750	NA	NA	NA	NA f	alse	NA :	AN		
5	524	NA	NA	NA	NA f	alse	NA :	AN		
6	619	259	649	NA	NA f	alse	NA :	NA		
7	312	743	619	NA	NA f	alse	NA :	AN		
8	440	1232	NA	NA	NA f	alse	NA :	NA		
9	352	978	827	NA	NA f	alse	NA :	AN		
10	714	NA	NA	NA	NA f	alse	NA :	AN		
11	781	922	NA	NA	NA f	alse	NA :	NA		

Figure 2: The figure shows the table format in the importing evidence stain file.

- Import population frequencies:

- o Requires an own folder (population-folder) with **only** frequency-files.
- o File-format:
 - Filename:
 - The name of the filenames **needs** to be on the form "kit_population.ext", where ext can be any extensions (or be missing as well).
 - kit="kit-name" and population="population name"
 - The kit-name must be consistent with the short-name of the kit instrument. See ?plotEPG for more details.
 - File:
 - First column needs to be allele-information (header-name may be anything).
 - Other columns are frequency-information (header-name denotes the locus name (loci names are converted to capital letters)).

- o To import frequencies:
 - Push "1) **Select directory**" button to select the population-folder with the population frequency files.
 - Push "2) **Import from directory**" button to import the population frequency files from the selected folder.
 - It is possible to add new files into the selected population-folder at any time and push the button once again to include new information to the dropdown-list.
- Selection of kit and population:
 - After importing the frequency-files (after pushed (2)), the user may select wanted kit and population from the two drop down lists at any time* (*not after a reference-database file has been imported).
 - This can be useful to see the EPG layout for different selected kits.
- Import Evidence/Reference sample (see figure 2 and figure 3):
 - o **Multiple** evidence or reference profiles are **allowed** in each file.
 - o In evidence files:
 - "height" header is required for analysis Deconvolution, Weight-of-Evidence (continuous model) and 'Database search'. For 'Qualitative LR' this is not required.
 - o In reference files:
 - "height" header is optional but will not be used further in any analysis.
 - o Note:
 - The import function will not check:
 - That the length of allele and heights are equal long for a given locus.
 - Loci without any allele-information (i.e. empty or dropped out), are NOT imported.

[1]	"Raw fil :	import:"		
	SampleName	Marker	Allele1	Allele2
1	Victim	D3S1358	16.0	15.0
2	Victim	TH01	9.3	9.3
3	Victim	D21S11	29.0	27.0
4	Victim	D18S51	17.0	15.0
5	Victim	D2S1338	23.0	19.0
6	Victim	D16S539	11.0	12.0
7	Victim	VWA	14.0	17.0
8	Victim	D8S1179	14.0	15.0
9	Victim	FGA	22.0	21.0
10	Victim	D19S433	13.0	15.2
11	Suspect	D3S1358	16.0	15.0
12	Suspect	TH01	6.0	7.0
13	Suspect	D21S11	29.0	35.0
14	Suspect	D18S51	11.0	14.0
15	Suspect	D2S1338	17.0	20.0
16	Suspect	D16S539	9.0	10.0
17	Suspect	VWA	15.0	17.0
18	Suspect	D8S1179	10.0	13.0
19	Suspect	FGA	22.0	25.0
20	Suspect	D19S433	14.0	14.0
•				

Figure 3: The figure shows the table format in the importing reference file.

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- **Import Reference Database** (see figure 4):
 - o Exactly same format as reference files.
 - o Multiple database file may be imported (**must** be done one-at-the-time).
 - o **Requires** that population frequencies are imported and selected.
 - WARNING: Population frequencies may not be changed again after database importing!
 - o Note:
 - The ranking of databases are done over all selected databases.
 - Same samples within a database needs to be in same block but markers within sample can be different orders.
 - Some samples may have more/less markers than others (e.g. SGMplus profiles contra ESX17).
 - **Missing markers** for a sample are given with NA.
 - Only markers shared with selected population frequencies are imported.
 - The imported database files may contain different markers.
 - Homozygote genotype may have an empty allele under 'Allele 2'.
 - The database file may contain any number of individuals.
 - o Tips:
 - It is more efficient to import several small databases than one big.
 - Time usage to import a database file with 16 markes:
 - o 1e6 profiles takes about 130 seconds
 - Requires ~1.3GB memory
 - o 5e6 profiles takes about 800 seconds.
 - Requires ~6.1GB memory
 - Save a lot of time and memory by storing a project to file (See File under toolbar). The imported database will be stored very efficiently.

```
[1] "Raw fil import:"
                           Sample.Name
                                         Marker Allele.1 Allele.2
   00-JP0001-14 20142342311 NO-3241 D3S1358
                                                       14
   00-JP0001-14_20142342311_NO-3241
                                         THO1
                                                                9.3
                                         D21S11
   00-JP0001-14_20142342311_NO-3241
                                                       29
                                                                 30
   00-JP0001-14_20142342311_NO-3241
                                         D18S51
                                                       13
                                                                 17
   00-JP0001-14_20142342311_NO-3241 D10S1248
   00-JP0001-14_20142342311_NO-3241 D1S1656
   00-JP0001-14 20142342311 NO-3241 D2S1338
   00-JP0001-14_20142342311_NO-3241_D165539
00-JP0001-14_20142342311_NO-3241_D22S1045
                                                       10
                                                       15
                                             VWA
10 00-JP0001-14_20142342311_NO-3241
                                                       17
                                        D8S1179
11 00-JP0001-14_20142342311_NO-3241
                                                       12
                                                                 13
12 00-JP0001-14_20142342311_NO-3241
                                             FGA
                                                       19
13 00-JP0001-14 20142342311 NO-3241
                                         D2S441
                                                       11
14 00-JP0001-14_20142342311_NO-3241 D125391
15 00-JP0001-14 20142342311 NO-3241
                                        D195433
16 00-JP0001-14 20142342311 NO-3241
                                            SE33
17 00-JP0001-14_20142342311_NO-3241 AMEL
18 00-JP0002-14_20142342311_NO-3242 D3S1358
                                                       15
                                                                 18
19 00-JP0002-14_20142342311_NO-3242
                                          TH01
20 00-JP0002-14_20142342311_NO-3242
                                         D21S11
                                                       28
21 00-JP0002-14_20142342311_NO-3242
                                         D18S51
22 00-JP0002-14_20142342311_NO-3242 D1051248
23 00-JP0002-14 20142342311 NO-3242 D1S1656
24 00-JP0002-14_20142342311_NO-3242 D2S1338
25 00-JP0002-14_20142342311_NO-3242 D16S539
                                                       25
                                                       11
                                                                 13
26 00-JP0002-14_20142342311_NO-3242 D22S1045
                                                       15
   00-JP0002-14 20142342311 NO-3242
                                                       14
```

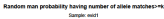
Figure 4: The figure shows the table format in the importing reference database file.

VIEW DATA:

- **View frequencies** (see figure 5 for the Norwegian SGMPlus population):
 - o Creates a new window which shows the selected population frequencies in a table.
 - o If any evidence profiles(s) are selected after evidence-import, the software makes a 'false positive probability' plot for each selected profiles.
 - The plot (figure 6) shows the probability that a random individual ('false positive probability') matching at least (2*n-wildcardsize) up to 2*n alleles (MAC) with a **selected evidence** profile. Here **n** is number of considered loci (which are both in evidence and population frequencies) and wildcardsize is number of allowed mismatches (default is wildcardsize =7).
 - wildcardsize can be changed under "Frequencies" in Toolbar by changing value **Set number of wildcards in false positive match.**
 - o Note:
 - Only allele-information in evidence-profiles are used.
 - New alleles which are not found in the selected population are assumed to have allele-frequency 0.

Allele	D3S1358	TH01	D21S11	D18S51
5	NA	0.00259844093543874	NA	NA
6	NA	0.209274435338797	NA	NA
7	NA	0.212472516490106	NA	0.000898472596585804
8	NA	0.0836498101139316	NA	NA
8.2	NA	NA	NA	NA
9	NA	0.140915450729562	NA	0.000998302885095338
9.3	NA	0.344293423945633	NA	NA
10	0.00089865202196705	0.00589646212272636	NA	0.0105820105820106
11	0.00559161258112831	0.000899460323805717	NA	0.00638913846461016
11.3	NA	NA	NA	NA
12	NA	NA	NA	0.132075471698113
13	0.00329505741387918	NA	NA	0.127882599580713
13.1	NA	NA	NA	NA
13.2	NA	NA	NA	NA
14	0.124113829256116	NA	NA	0.181291803933313
14.2	NA	NA	NA	NA
15	0.270993509735397	NA	NA	0.139862234201857
15.2	NA	NA	NA	NA .

Figure 5: The figure shows the viewed frequencies for the Norwegian SGMPlus frequencies.



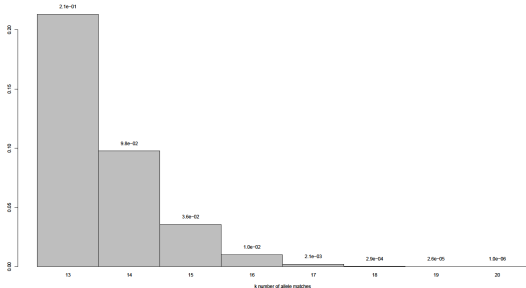


Figure 6: The figure shows the random match probability of matching with at least k number of alleles (in reference) with the observed alleles in evidence.

- **View evidence** (for selected evidence):
 - Prints imported alleles (and peak heights if any) for each selected evidence profile(s) (see figure 7).

```
[1] "Samplename: evid1"
       Allele Height
       "X/Y"
                     "2136/1015"
AMET.
D3S1358 "14/15/16"
                     "178/2405/1982"
       "6/7/9.3"
                     "419/282/1871"
TH01
D21S11 "27/29"
                     "1128/1750"
D18S51 "15/17"
                     "467/524"
D2S1338 "17/19/20/23" "290/619/259/649"
D16S539 "9/10/11/12" "217/312/743/619"
       "14/15/17" "1250/440/1232"
D8S1179 "10/13/14/15" "206/352/978/827"
        "21/22"
                     "664/714"
D19S433 "13/14/15.2" "1157/781/922"
```

Figure 7: The figure shows the printed alleles and heights in the imported evidence.

- o Plots EPG(s) (see figure 7) for each selected evidence profile(s)
 - Requires that user have imported "Population frequencies".
 - The kit selected under 'Select kit' denotes the EPG format.
 - Loci in evidence which are **inconsistent** with the ones in selected kit (or missing) are **not shown** in plot.
 - Evidence profiles without peak heights for corresponding alleles are given with peak height equal 1.
- o Note:
 - See ?plotEPG to see which kit-formats that are supported.
 - Reference profiles can be imported as evidence profiles and shown in a EPG.

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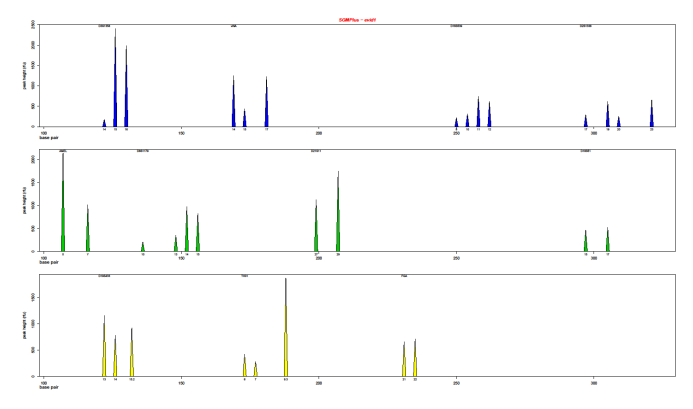


Figure 8: The figure shows the plotted EPG (on selected SGMPlus kit format) of the imported evidence stain.

- **View reference** (for selected reference):
 - o Prints imported genotypes for each selected reference profile(s) (figure 9).
 - o If any evidence profiles(s) are selected after evidence-import, the software counts number of matching alleles (MAC) for each loci of the selected reference profiles, for each selected evidences (figure 10).
 - MAC = number of alleles for the reference which are included in the evidence.
 - nLocs = number of considered loci when counting MAC.

```
Victim
                      Suspect
D3S1358 "16/15"
                      "16/15"
                      "6/7"
TH01
         "9.3/9.3"
         "29/27"
                      "29/35"
D21S11
         "17/15"
D18S51
                      "11/14"
D10S1248 "15/13"
                      "13/13"
D1S1656 "12/17.3"
D2S1338 "23/19"
D16S539 "11/12"
D22S1045 "15/16"
         "14/17"
D8S1179
        "14/15"
         "22/21"
D2S441
         "10/14"
D12S391
         "18.3/22"
                      "18/19"
D19S433
         "13/15.2"
                      "14/14"
         "30.2/33.2" "27.2/29.2"
```

Figure 9: The figure shows the printed alleles of the imported reference profiles.

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[1]	"Number	of	matching	alleles	with	samplename	evid1:	
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	Victim	Suspect
AMEL	NA	NA
D3S1358	2	2
TH01	2	2
D21S11	2	1
D18S51	2	0
D2S1338	2	2
D16S539	2	2
VWA	2	2
D8S1179	2	2
FGA	2	1
D19S433	2	2
MAC	20	16
nLocs	10	10

Figure 10: The figure shows number of matching alleles and total (MAC) with the imported and selected evidence stain. By combining the observed MAC and figure 7, the random match probability of observing MAC is useful for providing an extended version of "Random man not excluded"-statistics: The random match probability for Victim (MAC=20) becomes 1/1000000, while only 1/100 for Suspect (MAC=16).

- **View database** (see figure 11 for selected database):
 - Creates a new window (for each selected database) which shows the genotypes for every reference in the database.
 - "NA" means that the genotype of a reference was missing.
 - o If any evidence profiles(s) are selected after evidence-import, the software counts number of matching alleles (MAC) for all references in the database against each of the selected evidences (see figure 12). The results are shown in a MAC-ranked table in a new window (for each selected database).
 - MAC = total number of alleles for the reference which are included in the evidence.
 - Summed over all selected evidences.
 - **nLocs** is number of reference-loci which has been used to evaluate the MAC.
 - o Note:

Max number of individuals to view in a database can be changed with selecting
 Set maximum view-elements under "Database search" in toolbar.

Figure 11: The figure shows the viewed references inside the imported ESX17 database which are presented only with SGMPlus profiles since the selected kit for the imported frequencies was SGMPlus_Norway.

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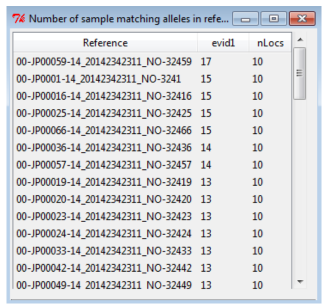


Figure 12: The figure shows the sorted references (in the reference database) with respect to MAC (total number of matching alleles) to the selected evidence.

specified hypothesis using the continuous model as described in the vignette. 333 Requires: Imported population frequencies. 334 335 o Feature: Allele drop-out, Drop-in (with a peak height model) and stutter. 336 337 **Deconvolution:** 338 Deconvolution ranks the most probable combined genotype profiles given a specified **hypothesis** and the Maximum Likelihood Estimates of the parameters in the continuous 339 model (as given in the vignette). 340 Requires: Imported population frequencies and selection of at least one evidence profile 341 with peak height information. References are optional to condition on in the hypothesis. 342 Feature: Model may handle replicates, allele drop-in, drop-out and stutter. 343 344 345 Weight-of-Evidence: Weight-of-Evidence is done by comparing the Likelihood Ratio (LR) between the 346 specified hypotheses Hp (prosecution) and Hd (defence) using the continuous model as 347 given in the vignette. 348 Modules: 349 350 1) 'Continuous LR' (Maximum Likelihood based) Optimizes (maximum) the model parameters in the continuous model. 351 2) 'Continuous LR' (Integrated Likelihood based) 352 353 Integrates out the model-parameters in the continuous model. 3) 'Qualitative LR' (semi-continous) 354 355 Explores LR as a function of allele dropout probability parameter. 356 357 Requires: 358 Imported population frequencies, at least one evidence profile and at least one 359 reference profile (suspect) to weight evidence for. Additional reference profiles are optional to condition on in the hypotheses. 360 'Continuous LR' requires evidence(s) including peak heights, 'Qualitative LR' 361 only requires allele data. 362 363 Feature: 364 The continuous model: Handles replicates, allele drop-in, drop-out, stutter and 365 fst-correction. 366 The semi-continuous model: Handles replicates, allele drop-in, drop-out and fstcorrection. 367 368 369 370 **Database search:** o Does weight-of-evidence by comparing the Likelihood Ratio (LR) between the specified 371 372 hypotheses H_i (reference i in database) and Hd (defence) using the continuous model as given in the vignette. 373 o Modules: 374

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Generate sample:

INTERPRETATIONS:

o Generates alleles using the population frequencies and draws peak heights for a

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- 1) 'Continuous LR' (Maximum Likelihood based)
- 2) 'Continuous LR' (Integrated Likelihood based)
- 3) 'Qualitatitve LR' (Semi-continuous based)
- o Requires: Imported population frequencies, **at least one** evidence profile with **peak height** information and **at least one** reference-database. Reference profiles are optional to condition on in the hypotheses.
- o Feature: Model may handle replicates, allele drop-in, drop-out, stutter and fst-correction.
- o The continuous LR value is showed together with qualitative LR and MAC.

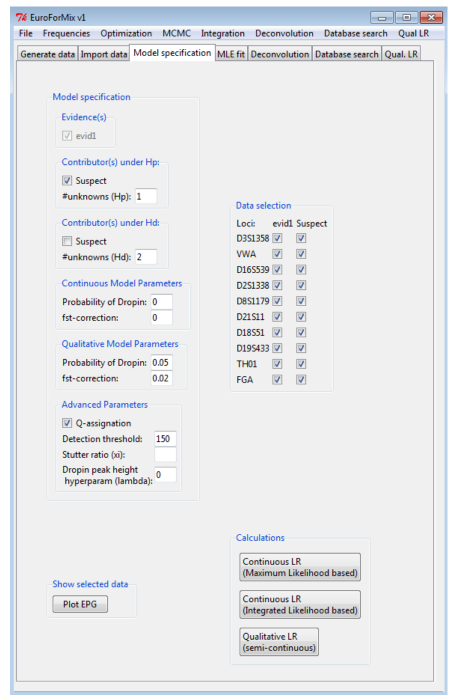


Figure 13: The figure shows the <u>Model Specification</u> GUI page for **Weight-of-Evidence** based on Likelihood Ratio calculation.

1		MODEL SPECIFICATION
23	- Evide	nce(s):
4	0	Shows selected evidence(s) from 'Import data'.
.5	0	All interpretations support multiple replicates.
5 7		 Note: All replicates are assumed to have same parameter sets.
	- Contr	ributors under Hp
	0	Case: Weight-of-Evidence or 'Database search'):
		 User may condition on selected references (from 'Import data') in the hypothesis
		Hp.
		 #unknowns under Hp: Denotes number of unknown contributors under the
		prosecution hypothesis Hp.
	0	Case: 'Database search':
		 The individual in the reference-database is already included in the hypothesis
		Hp.
	0	Case: Deconvolution or 'Generate sample':
		This block is not considered, since Deconvolution only considers the model
		under Hd, and sample generation is done only under a specific hypothesis.
	- Contr	ributors under Hd (same for all cases):
	0	User may condition on selected references (from 'Import data') in the hypothesis Hd.
	0	#unknowns under Hd: Denotes number of unknown contributors under the prosecution
		hypothesis Hd.
	0	Case: Weight-of-Evidence or 'Database search':
		References which are conditioned under Hp but not under Hd, will be assumed
		to be a ' known non-contributor ' under Hd (this is relevant when fst>0).
	- Conti	nuous Model Parameters and Qualitative Model Parameters:
	0	The Continuous Model Parameter section is only used for "Continuous LR"
		Calculations, while Qualitative Model Parameters section is only used for 'Qualitative
		LR' Calculations.
	0	'Probability of drop-in': [0,1]
		• Assumed probability of a random allele drop-in to the evidence at a given locus.
		See vignette for more details.
		This is default 0 for continuous models and 0.05 for qualitative models.
	0	fst-correction: [0,1]
		 Assumed co-ancestry parameter assigned in the genotype probability for each contributor in the hypotheses. See vignette for more details.
		• • • • • • • • • • • • • • • • • • • •
	_	 This is default 0 for continuous models and 0.02 for qualitative models. Case 'Database search':
	0	
		 When doing database search with "Continuous LR" Calculations, the allele drop- in probability for the qualitative LR can be changed by Set drop-in probability
		for qualitative model under "Database search" in toolbar (default is 0.05).
		When doing database search with "Qualitative LR" Calculations, this value is
		ignored in favor of the specification under "Qualitative Model Parameters".
		ignored in lavor of the specification under Quantative Model I arameters.

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470	- Adva	nced Parameters
471	0	Q-assignation:
472		• If checked, all alleles not present in the evidence are considered as allele "99".
473		Its frequency will be given as the sum of the frequencies for all the "non-
474		present" alleles.
475		If unchecked, the original alleles in the population are used as before.
476	0	'Detection threshold': [0,->)
477		 The threshold of required allele peak heights of whether an allele is present in
478		the evidence or not.
479		 Note: If peak heights in evidence are lower than the specified threshold,
480		the corresponding alleles (and peak heights) below threshold are
481		removed automatically. This may cause some loci to become empty.
482	0	'Stutter ratio': [0,1]
483		 Only used for 'Continuous LR' Calculations.
484		• Stutter ratio is a constant parameter "xi" which denotes the proportion of peak
485		heights from allele 'a' which is added to allele 'a-1'. See vignette for more
486		details.
487		• If allele 22 with peak height y_22 is contributed by a contributor and
488		allele 23 did not have any observed peak height, then the stutter
489		contribution to allele 21 from allele 22 will be (xi * y_22).
490	0	'Dropin peak height hyperparam': [0,1]
491		 Only used for 'Continuous LR'.
492		 Assumed hyper-parameter to model the peak height of the dropped in allele
493		caused by a 'random allele drop-in' if 'Probability of drop-in'>0. See vignette
494		for more details.
495		
496	- 'Data	base(s) to search' (case: 'Database search')
497	0	Lists the selected imported reference-database(s) to do the database search for.
498		
499		
500		DATA SELECTION
501		
502	- Select	t/unselect loci:
503	0	The user may select or unselect loci for each selected evidence(s) and reference(s) from
504		"Import data"
505	0	If a locus has been unselected for any of the evidence(s) or reference(s), the unselected
506		locus will not be evaluated at all.
507	0	Note: Evidence with more than 30 loci will not be able to be selected.
508		
509	- Missi	ng data:
510	0	Data with missing allele in any of the loci will automatically be deselected (inactivated)
511		such that the corresponding loci will be unavailable to evaluate.
512	0	For continuous I R evaluation:

• The Qualitative Model Parameters section is removed.

o Case **Generation** and **Deconvolution**:

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• If peak heights (in any of the evidence(s)) are missing for any selected locus, the user gets a message about deselecting the issued loci before proceeding.

- New alleles:

o If new alleles (does not exist in the population frequency table) occurs in the imported evidence or reference profile, the new alleles are assigned with allele frequency 'freq0'. 'freq0' is equal minimum observed frequency in population if N=0, or 'freq0'=5/(2N) where N is size of imported frequency database under "Frequencies" in Toolbar. The frequencies are after normalized.

SHOW SELECTED DATA

- Plot EPG:

- o **Prints** the selected evidence sample(s), reference(s) and considered population frequencies which are eventually used for further analysis **out to terminal**.
- o The selected evidence samples are shown in an EPG-plot.
 - Note: Alleles with corresponding peak heights below the specified "Detection Threshold" are removed.

CALCULATIONS

- 'Continuous LR (Maximum Likelihood based) ' (case Weight-of-Evidence and 'Database search'):
 - Maximizes the Likelihood of the unknown parameters in the continuous model given the assumed model so they attain maximum values for the specified hypothesis Hd (and Hp in case of Weight-of-Evidence).
 - The optimizer should return a global maximum. However, it may sometimes just return a local maximum. Number of start-points should be increased to ensure that the optimizer finds the global maximum of the Likelihood function. This can be changed under "Optimization" in Toolbar.
 - o After calculation, the page 'MLE fit' is visited to present maximized results.
- 'Continuous LR (Integrated Likelihood based)' (case Weight-of-Evidence and 'Database search'):
 - o Instead of optimizing the Likelihood of the unknown parameters, a **multivariate integration** over the unknown parameters are applied both under hypothesis Hp and Hd.
 - The accuracy of the integral depends on the specified 'relative error requirement' (see vignette for details).
 - Can be changed under "Integration" in Toolbar. Default is 0.005.
 - o In the output (see Figure 14), also the relative error of the LR is given in brackets.
 - o The integral requires that an **upper boundary** for the parameters mu (amount of DNA) and sigma (coefficient of variation) are specified. As default these are 20000 and 1, respectively. These values may be changed under "Integration" in Toolbar. See vignette for details.
 - o Calculates LR-values directly and avoids visiting the tab 'MLE fit'.

- Case Weight-of-Evidence: A message with LR pops up after calculation (see Figure 14).
- Case 'Database search': Database search results are shown directly after calculation (goes to tab 'Database search').
- 'Continuous LR (Integrated Likelihood based)' is not possible for multiple replicates and large number of loci since it doesn't evaluate on log-scale. Use the Maximum Likelihood based method instead if the other method goes wrong.

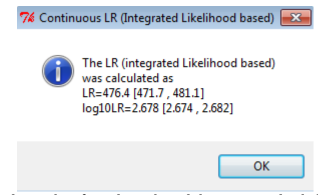


Figure 14: The figure shows the calculated Weight-of-Evidence based the Integrated Likelihood based continuous LR for the specified model in Figure 13.

- 'Qualitative LR (semi-continuous)' (case Weight-of-Evidence)

- Performs a semi-continuous procedure where the distribution of the 'allele drop-out probability given number of observed alleles' are utilized to infer a "conservative" LR.
 - The model is purely qualitative which means it is only based on alleleinformation.
- Goes directly to page Qual. LR.

- 'Generate sample' (case 'Generate sample'):

- A dataset (evidence sample and contributing references) will be randomly simulated under the specified model under "Model specification".
- o Reference profiles may be imported and selected as assumed known in the hypothesis.
- o Detection threshold, stutter ratio, probability of drop-in and drop-in peak height hyperparam may all be used in the simulation (**fst** are not used).
- The unknown contributor profiles under the hypothesis will be randomly generated using the selected population frequencies.
- The simulated peak heights of the evidence in the dataset are entirely based on the continuous model for assumed values of the model-parameters (mu,sigma,xi,mx).
 Default these are given as mu=1000, sigma=0.15, xi=0.1, mx=(C:1)/sum(C:1), where C is number of contributors.
- o Goes directly to page Generate data.

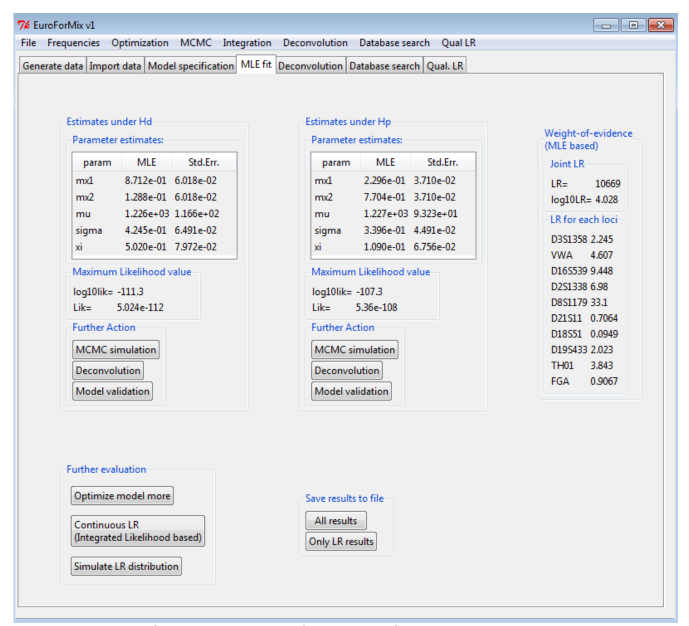
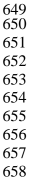


Figure 15: The figure shows the MLE-fit GUI page after doing **continuous LR** (**Maximum Likelihood based**) calculation (maximizing the continuous model with respect to the unknown parameters for each of the specified hypothesis in figure 13) for **Weight-of-Evidence**.

-	Parameter estimates:
	 param: The unknown parameters in the model (see vignette for more details). mx_i: Mixture-proportion for contributor 'i'. mu: Expected amount of DNA. sigma: Coefficient of variation. xi: Stutter ratio (fraction of peak height that are stutter).
	 MLE: The optimized¹ parameters in the model which attains a maximum point of likelihood function.
	 Std.Err.: The standard error of the parameter estimates in the model (see vignette f details).
-	Maximum Likelihood value:
	o log10lik and Lik: The ten-logged and the original value of the Likelihood value at from the optimization ¹ .
_	Further Action:
	o MCMC simulation (see Figure 16):
	 Performs 'Markov Chain Monte Carlo (MCMC) random walk Metropolis'
	samples under the desired hypothesis.
	 Uses the mode and the covariance matrix attained from the optimiz
	See vignette for details.
	• The first column in the output shows the estimated posterior distributions
	each of the unknown parameters in the model.
	• The second column in the output monitors the parameter samples in the
	simulation. • After sampling, the acceptance rate of the sampler is printed out to the term.
	The sampling, the acceptance rate of the sampler is printed out to the ter
	 Acceptance ratio = number of accepted samples divided by number proposed samples.
	 Ideally the acceptance rate should be around 0.2 to ensure that the
	parameter space has been fully explored.
	Tweak 'variance of randomizer' under MCMC in toolbar
	change the acceptance rate.
	 User may change number of required samples in the simulation under
	'MCMC' in toolbar.
	 The purpose of the MCMC simulation is to use it as an exploratory tool t
	 That the optimizer has found the global maximum.
	• The shape of the posterior distribution of the parameters.

¹ This may be only a local maximum point, not the global maximum (i.e. the Maximum Likelihood Estimate). Increase **number of start points** under "Optimization" in Toolbar to ensure a global maximum.



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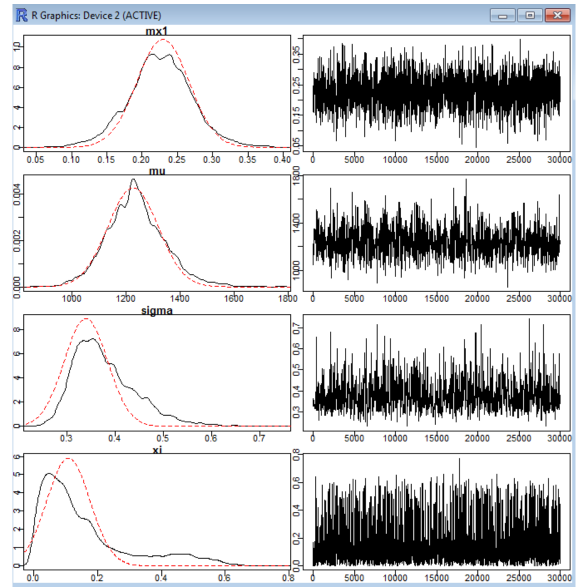


Figure 16: The figure shows the posterior density of the unknown parameters (first column) and corresponding iteration values (second column) from the MCMC method under the hypothesis Hp: "Suspect+1 unknown individual contributes to evidence evid1". The acceptance ratio was given as 0.35.

o **Deconvolution**:

Performs "Deconvolution" under the desired hypothesis. (See <u>Deconvolution</u> (page 5) for details.

Model validation (Figure 17):

- Uses a statistical hypothesis test to reject whether the maximum likelihood fitted model fits the observed peak heights (i.e. whether the gamma model assumption is reasonable).
- Estimates the cumulative probability of the observed peak heights conditional on the other peak heights (see vignette for more details).

- Uses a one-sample Kolmogorov-Smirnov test to test if the observed cumulative probability deviates significant from the uniform distribution.
- P-value from the test is printed out to terminal.
- A textbox is shown when the P-value is lower than the significance level 0.05 (i.e. rejection of assumption).

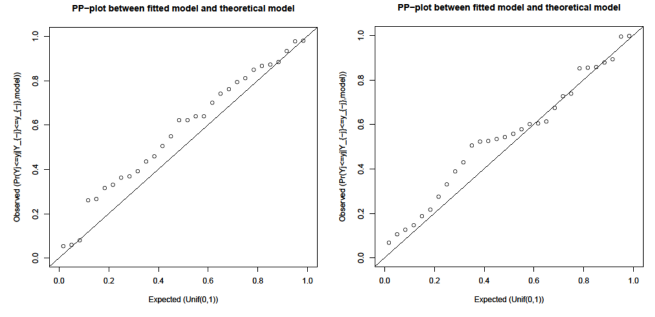


Figure 17: Left subplot shows the "**Model validation**" under hd with p-value 0.37. Right subplot is "**Model validation**" under hp with p-value 0.29.

WEIGHT-OF-EVIDENCE (case Weight-of-Evidence)

- Description:

- o The Weight-of-Evidence value is the ratio between the likelihoods of the two specified hypotheses Hp and Hd as specified in "Model specification".
- o The Weight-of-Evidence value is based on the continuous model as described in the vignette and handles allele drop-in, drop-out and stutter.

- Join LR:

- LR: 'Likelihood value under optimization under Hp' divided by 'Likelihood value under optimization under Hd'
- o log10: The ten-logged value of LR.

- LR for each loci:

- The LR for each loci separately (given the parameter-modes under Hp and Hd). See vignette for details.
- O Note: This will not be shown for evaluation of more than 30 loci

FURTHER EVALUATION

- Optimize model more:

- The optimization procedure can be run again with the same specifications as selected in "Model specification" to ensure that a global maximum is attained.
- o It is recommended to do this and check that the optimized Likelihood value is not increased further.
- Database search (case: 'Database search'):
 - o A database search with the specified continuous model will be applied. (See <u>Database</u> search for details.
- 'Continuous LR (Integrated Likelihood based)' (case Weight-of-Evidence)
 - o See CALCULATIONS under section "Model specification".
- 'Simulate LR distribution' (case Weight-of-Evidence)
 - o MCMC simulation will be applied both under Hp and Hd to provide a plot of a "Bayesian" distribution of the LR where the uncertainty of the parameters in the continuous model under both Hp and Hd are taken into account (see Figure 18).
 - Number of samples can be changed with Set number of samples under MCMC in Toolbar (default is 10000 samples).

Distribution of LR over posterior space of parameters

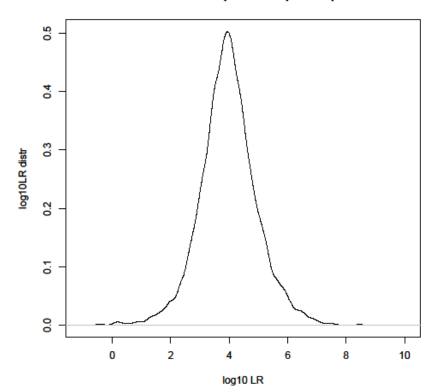


Figure 18: The plot shows the distributed LR where the *a posteriori* density of the parameters in the continuous model under both Hp and Hd are taken into account. *a posteriori* density are simulated using the **MCMC simulation** (Figure 16 shows only Hp).

- 'All results':

722 723 The parameter estimates with corresponding standard deviation errors estimates and the likelihood values will be printed to file for all hypotheses on page (see below).

param-MLE-Std.Err. mx1-0.87124-0.06018 mx2-0.12876-0.06018 mu-1226.3- 116.6 sigma-0.42447-0.06491 xi-0.50195-0.07972 log10Lik=-111.3 Lik=5.024e-112 -----Estimates under Hp----param-MLE-Std.Err. mx1-0.2296-0.0371 mx2-0.7704-0.0371 mu-1226.65- 93.23 sigma-0.33957-0.04491 xi-0.10902-0.06756 log10Lik=-107.3

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- 'Only LR results': (case Weight-of-Evidence)

Lik=5.36e-108

• The LR calculated values shown in WEIGHT-OF-EVIDENCE will be printed to file (see below).

Marker	LR	log10LR
D3S1358	2.245e+00	0.35113
VWA	4.607e+00	0.66345
D16S539	9.449e+00	0.97536
D2S1338	6.980e+00	0.84384
D8S1179	3.310e+01	1.51979
D21S11	7.064e-01	-0.15094
D18S51	9.490e-02	-1.02273
D19S433	2.023e+00	0.30610
TH01	3.843e+00	0.58467
FGA	9.067e-01	-0.04253
JointMLE	1.067e+04	4.02814

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4. Deconvolution:



Figure 19: The figure shows the <u>Model Specification</u> GUI page for doing **Deconvolution**. We condition on the suspect, and assume one unknown in the hypothesis. Our model assumes unknown "(n-1)- stutter" ratio, no allele drop-in and no theta-correction.

- Description:

- o Deconvolution is applied for a specific hypothesis Hd as shown in Figure 19.
- The deconvolution conditions on the optimized parameters (i.e. the <u>MLE fit</u> in Figure 20) for the continuous model.
- The deconvolution result shows (see Figure 21) a ranked list of the **posterior probabilities** of the combined genotype-profiles (see vignette for details).
- O Since the deconvolution is based on the continuous model it may handle multiple replicates, allele drop-in, drop-out and stutter.

- Table:

- The columns in the table (see Figure 21) show the resolved genotype for each contributor in the specified hypothesis (per locus).
- o The combined profiles are ranked due to their **posterior probabilities**.

- The ranked elements in the table ensures that the sum of the **posterior probabilities** are at least 0.9999.
 - Can be changed under 'Deconvolution' in toolbar.
- Maximum length of table is default 10000.
 - Can be changed under 'Deconvolution' in toolbar.
- o Note:
 - Having only sub-optimized parameters (in the MLE fit) will not give the most likely genotypes.
 - Q-assignation is recommended to use since dropped out alleles are equally threated and assigned as "99".

- Save table:

• The **full** table will be exported to a tabulator-separated text-file.

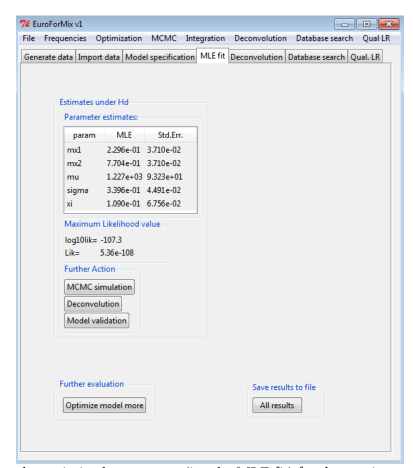


Figure 20: The figure shows the optimized parameters (i.e. the <u>MLE fit)</u> for the continuous model. The fitted model has the same "Further Action" possibilities as for "Weight-of-Evidence" and "Database search".

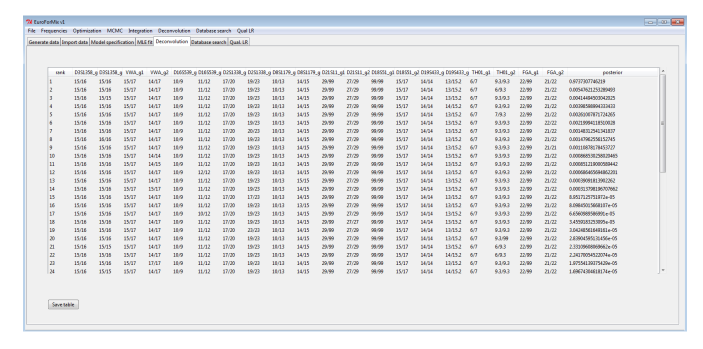


Figure 21: The figure shows the ranked table of deconvoluted genotype profiles for the unknown major contributor, when conditioning on the suspect profile. The table is ranked with respect to the posterior probability of different combined genotype profiles. The top ranked combined genotype profile is an outlier from the others which indicates that it is possible to extract the unknown profile (from figure 9 we see that this is a correct extraction).

5. Database search:

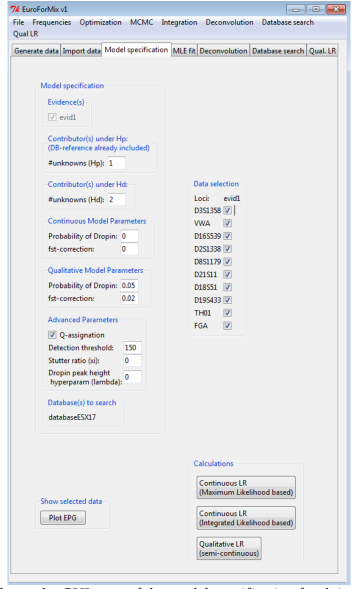


Figure 22: The figure shows the GUI page of the model specification for doing database search on the database file "databaseESX17". Our model assumes no "(n-1)-stutter", no allele drop-in and no theta-correction.

- Description:

The 'Database search' is very similar as the Weight-of-Evidence (see Figure 22) with the only difference in that each individual in the reference-database is assumed as a contributor in the hypothesis Hp. For each individual 'j' in reference-database we calculate a LR-value LRj.

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- The user may choose between using peak heights in a 'Continuous LR' (Maximum **Likelihood based** or **Integrated Likelihood based**)' calculation or ignoring the peak heights in a 'Qualitative LR' calculation.
- When selecting 'Continuous LR':
 - 'Qualitative LR' is always calculated along with the 'Continuous LR' values.
 - The qualitative model assumes an allele drop-out parameter which is estimated.
 - The allele drop-in parameter in the qualitative model is set as default 0.05, but can be changed with "Set drop-in probability for qualitative model" under 'Database search' in the Toolbar.
 - No theta-correction is assumed in the qualitative model.
 - If "Continuous LR (Maximum Likelihood based)" calculation is used, the optimized parameters under the Hd -hypothesis are first shown (see Figure 23).

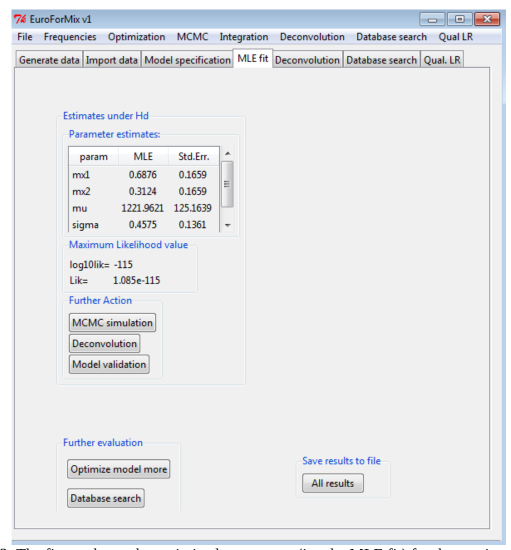


Figure 23: The figure shows the optimized parameters (i.e. the MLE fit) for the continuous model under Hd (with specifications as given in Figure 22). The fitted model has the same "Further Action"

839 possibilities as for "Weight-of-Evidence" and "Deconvolution". The user must push "Database search" for doing the actual database searching. 840 841 842 When selecting 'Qualitative LR': o The "Set drop-in probability for qualitative model" under 'Database search' in the 843 Toolbar is ignored. 844 845 o The qualitative model assumes an allele drop-out parameter which is estimated. 846 The 'Continuous LR' calculation is ignored. 847 848 Note: 849 The 'Continuous LR' calculation is based on the **continuous model** as given in the vignette and hence may handle allele drop-in, drop-out and stutter. 850 Continuous LR (Integrated Likelihood based) is not possible to use for replicates. 851 852 The reason for showing the MLE fitted parameters under Hd (see Figure 23) for "Continuous LR (Maximum Likelihood based)" calculation is that the user should have 853 854 the possibility to check if the parameter estimates under Hd seems reasonable so he can go back and change the model specification. 855 856 857 **Table** (see Figure 24): 858 859 860 861

- 'Reference name' is name of individuals given in the reference-database.
- The table shows the ranked individuals in the database due to the continuous LR values (contLR), qualitative LR values (qualLR), number of matching alleles (MAC) or number of evaluating loci (nLocs).
- qual.LR (Qualitative LR (semi-continuous model))
 - Parameter for dropout probability is based on the median of 2000 samples from the 'distribution of dropout-probability'.
 - Number of required samples may be changed under 'Qual LR' in toolbar.
 - For multiple evidences, the mean of the median is used as the dropout probability parameter.
 - Assumes drop-in probability 0.05 as default. Can be changed under 'Database search' in toolbar.
 - Assumes no theta-correction.
- o MAC (Matching allele counter) is number of alleles in the reference-profile which matches the evidence.
 - Note: MAC is summed over the considered evidences.
- o **nLocs** is number of loci in the reference-profile which are used to calculate the contLR, qualLR and MAC.
 - Note: Some references in the database may be missing loci which are presented in the evaluated evidence.

Note:

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- Maximum number of elements to view a 'Database search' result table is 10000. This can be changed under 'Database search' in toolbar.
- Putting fst>0 may be very time-consuming since we require that individual 'j' is a known non-contributor under Hd, and hence Hd is calculated for each individual in database.

- If no allele drop-in is assumed under the continuous model, **cont.LR** is not calculated for the non-fitting individuals in the database.
- Save table:
 - The full table will be exported to a tabulator-separated text-file.

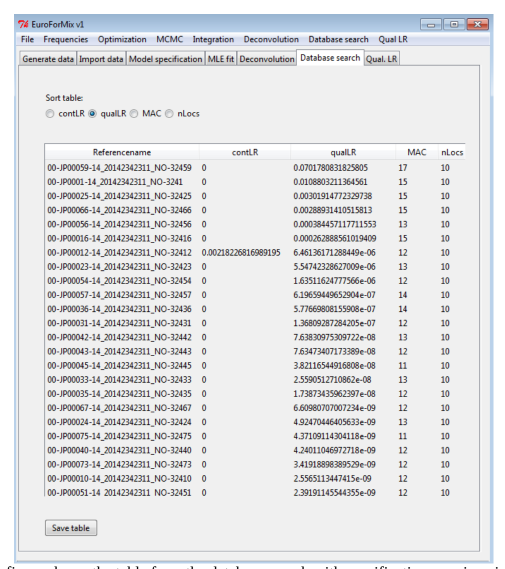


Figure 24: The figure shows the table from the database search with specifications as given in Figure 22 based on 'Continuous LR' (Maximum Likelihood based)" calculations. The references are sorted due to the qualitative LR's (which assumes allele drop-out probability 0.08 and allele drop-in probability 0.05).

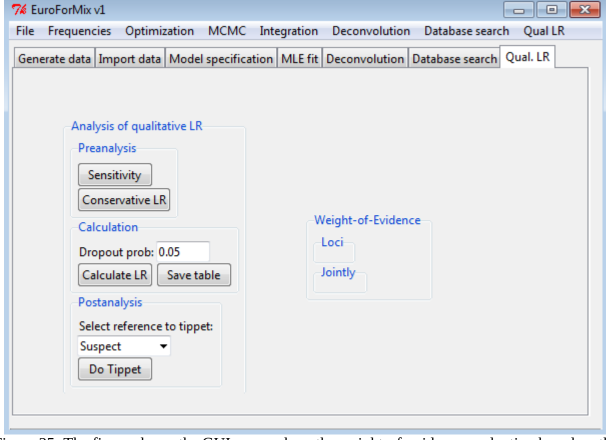


Figure 25: The figure shows the GUI page where the weight-of-evidence evaluation based on the qualitative model is done.

- Description:

O This module samples from the distribution of the 'allele drop-out probability given number of observed alleles' to evaluate the qualitative LR automatically. Also sensitivity plot as a function of allele-dropout probability and random man tippet analysis is implemented.

PREANALYSIS

- Sensitivity:

- o Plots the log10LR as a function of allele-dropout probability (see Figure 26).
 - The upper probability range and number of ticks can be changed under 'Qual LR' in the toolbar.
- o Note:
 - Lower range in sensitivity is 1e-6 (something small).

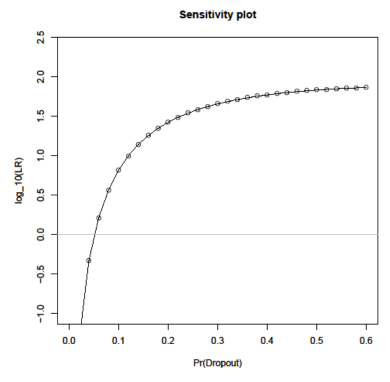


Figure 26: The figure shows the plot of Weight-of-evidence (Likelihood Ratio) as a function of allele drop-out probability.

- Conservative LR:

- O By sampling from the "allele drop-out probability given number of observed alleles in the evidence"- distribution for the hypothesis Hp and Hd, the most 'conservative' LR (i.e. smallest) is automatically calculated and printed (see Figure 27 and Figure 28).
 - The most "conservative" LR is found by following:
 - Take out the "alpha" and "1-alpha"-quantiles from the simulated 'allele-dropout probability distribution' under both Hp and Hd.
 - The quantile (under both Hp and Hd) which gives the lowest LR is the "conservative LR".
 - The significance level "alpha" is given 0.05 as default.
 - This can be changed under 'Qual LR' in the toolbar.
 - The number of required samples from the 'allele-dropout probability distribution' is given 2000 as default.
 - This can be changed under 'Qual LR' in the toolbar.
 - Note: If no samples are accepted from the allele-dropout probability distribution', an error-message is provided to the user.
- When more evidence samples are imported, the most 'conservative LR' over all samples is considered.
 - The dropout probability quantiles are estimated for each of the evidence samples.

Figure 27: The plot shows the sampled 5% and 95% quantiles of the distribution of the 'allele drop-out probability given number of observed alleles'.

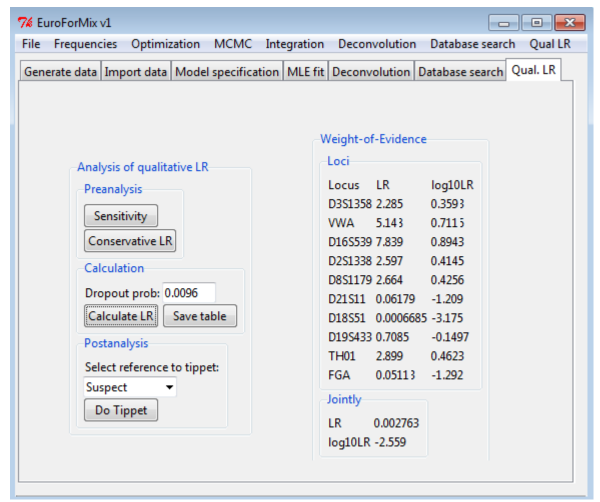


Figure 28: The plot shows the conservative Weight-of-Evidence values (Likelihood Ratios) after pushing "Conservative LR". The most conservative estimated allele drop-out probability-quantile from Figure 27 was the 5% quantile under Hd which gave 0.0096. Hence the table in this plot shows the LR inserted for this value.

963 CALCULATION 964 965 **Dropout prob:** 966 • The user may specify the assumed number of allele dropout-probability. 967 Calculate LR Instantly calculates the LR for the given user-specified allele dropout probability in 968 969 "Dropout prob". 970 Save table: 971 o Saves the weight-of-evidence calculated LR results to a selected file. 972 973 974 **POSTANALYSIS** 975 976 **Selection of reference to tippet:** 977 A drop-down list of references which are conditioned under Hp but not under Hd. 978 979 **Do Tippet:** 980 Random tippet samples are provided by replacing the selected reference (under the dropdown list in the hypothesis Hp) with a random individual from the population and then 981 calculate his LR. A vast amount (default is 1e6) of random tippets is simulated to 982 determine the tippet-distribution. 983 984 The mean, standard errors of LR and log10LR-quantiles (1%, 5%, 50%, 95%, 985 99%) are printed out to terminal (see Figure 29). A plot of the cumulative distribution of log10LR will be shown (see Figure 30). 986 987 Number of tippets can be changed under 'Qual LR' in the toolbar. 988 If weight-of-evidence has been calculated: The reporting LR for the "tipped individual" is superimposed as a blue line to the 989 plot (see Figure 30). 990 991 The discriminatory metric (log10LR-q99%) is printed out to terminal (see Figure 992 993 Note: Precalculations are always done previous to the tippet-sampling, therefore the 994 number of tippets are only limited to make the plot. 995 996 [1] "Precalculating for tippet plot..."

997 998

999

1000 1001

1002

1003

```
[1] "Precalculating for tippet plot..."
[1] "Simulating 1e+06 tippets..."
[1] "Mean of samples = 0.703689922273713"
[1] "Standard Error of samples = 0.677592812037152"

1% 5% 50% 95% 99%

-32.630258 -28.495191 -18.700169 -9.622545 -6.242242
[1] "Discriminatory metric (log10(LR) - q99) = 3.6836388776166"
```

Figure 29: The plot shows the printed tippet information to the terminal when replacing the "Suspect" in hypothesis Hp with a random man (a tippet). Number of tippets simulated, mean and standard errors of LR and log10LR-quantiles (1%, 5%, 50%, 95%, 99%) are printed out to terminal (see Figure 29). Also the discriminatory metric, the distance between the observed log10LR for the suspect and log10LR-99%-tippet-quantile is given.

Tippet calculation for Suspect with 1e+06 samples.

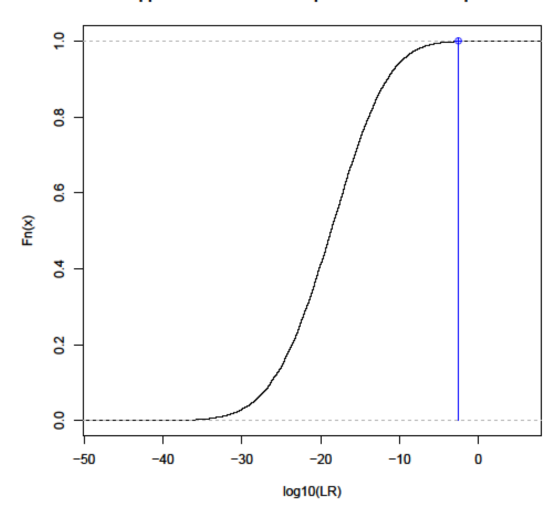


Figure 30: The figure shows a cumulative distribution of 1000000 log10LR tippets, where each tippet is based on replacing the "Suspect" in hypothesis Hp with a random man from the population. The reporting LR for the "tipped individual" (i.e. "Suspect in this case) is superimposed as a blue line to the plot.

 $\frac{1004}{1005}$

 $\begin{array}{c} 1007 \\ 1008 \end{array}$

7. Generate data:

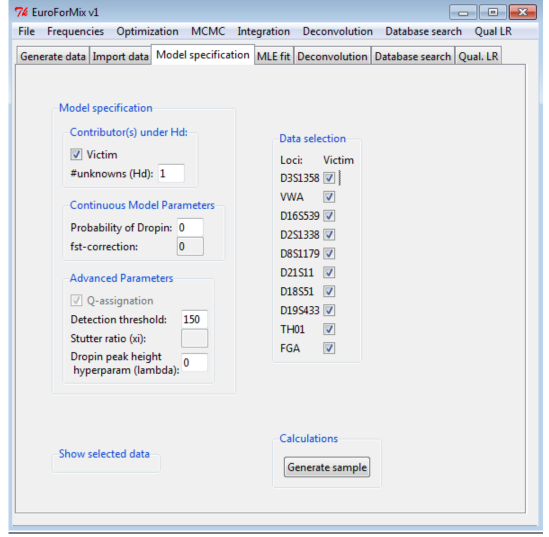


Figure 31: The figure shows the <u>Model specification</u> GUI page for generating allele with corresponding peak heights from the continuous model for a given specified model. From here we will generate data which are contributed from a known Victim profile and an unknown individual. We assume a detection threshold of 150 rfu and no allele drop-in is considered.

- Description:

 Generates alleles using the population frequencies and simulates peak heights for a specified hypothesis (see Figure 31) using the continuous model.

 The generation may simulate allele-dropout, drop-in (with a peak height model) and stutter (see Figure 32).

Allele-dropout is indirectly simulated by falling below the defined threshold.

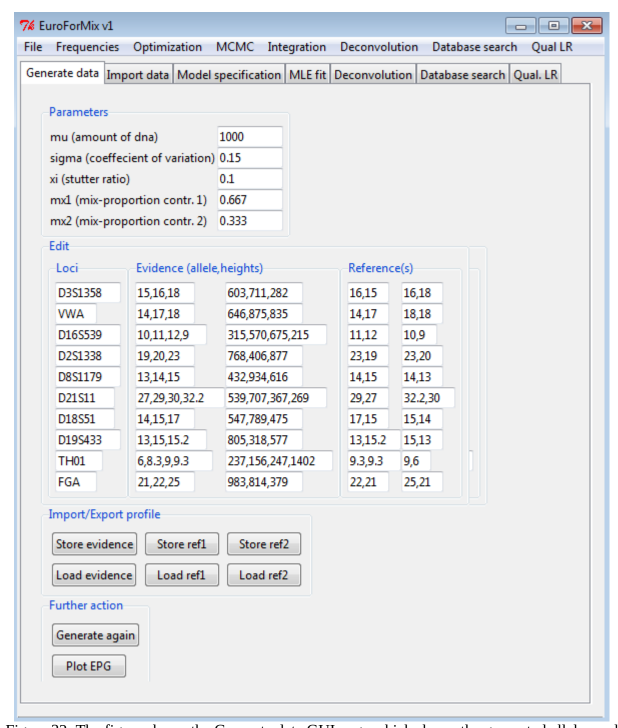


Figure 32: The figure shows the <u>Generate data</u> GUI page which shows the generated alleles and corresponding peak heights (under **Evidence**) for the given selected set of parameters under **Parameters**. The true contributors are given under **Reference**(s).

1043 1044 **Parameters:** 1045 o **mu**: amount of DNA 1046 o **sigma:** coefficient of variance o **xi:** stutter ratio 1047 mx=(mx1,..., mxC): mixture proportion for contributor 1,...,C. 1048 1049 • Note: **mx** will be normalized if it's not already. 1050 1051 Edit: 1052 o **Loci**: Loci name of the population frequency used to generate the dataset. **Evidence**: The allele information is given in the left column while the peak height 1053 information is given in the right column. Each element **needs to be** separated with ",". 1054 **Reference**: The alleles of the true contributors to the generate evidence is sequentially 1055 shown in each column. 1056 All the loci names, evidence-allele and heights and reference-alleles may be edited 1057 1058 before storing (See Figure 32). 1059 Import/Export: 1060 o Save data: 1061 1062 Stores the generated (and possible edited) evidence- or reference-profile to a file. Extension .csv added automatically. 1063 Load data: 1064 Loads profiles from file into the selected entries (evidence or reference). 1065 This is useful for generating random evidence samples where loaded 1066 references are conditioned on. 1067 1068 Note: 1069 If any locus is missing from the loaded evidence or reference file, the edit-cell will be empty. 1070 The order of the loci in the file does not matter. 1071 1072 1073 **Further action:** 1074 Generate again: Make a new simulation of the evidence sample using the selected values of the parameters under **Parameters**. 1075 Plot EPG: Plots the generated (and possible edited) evidence in a EPG-plot. 1076 It will use the "kit" selected under "Import Data"-page. 1077 See ?plotEPG to see which kit-formats that are supported in the EPG. 1078 1079 1080 1081 1082 1083 1084 1085 1086 1087 1088

(C) To be implemented in a future version:

1090

- Label the alleles of the selected references to the EPG-plot.
- Warning if exp(lik)=0 when lik>-Inf (happens for INT calculations)
- Empty loci will not be removed when imported to the software. They will be considered as a full dropped out loci in the evaluation.