LRmix Studio 2.1.1 user manual

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1. What is LRmix Studio?

LRmix Studio is a free of charge, open-source (GPLv 3 license), expert system dedicated to the interpretation of forensic DNA profiles, with a particular focus on complex DNA mixtures. LRmix Studio enables measuring the probative value of any (autosomal STR-based) forensic DNA profile.

LRmix Studio is programmed after the likelihood ratio model described in Haned et al (FSIG 2012) and Gill & Haned (FSIG 2013). This model explicitly accommodates for uncertainty in the DNA profiles from the allelic drop-out and drop-in phenomena. The program estimates these quantities from the available data, and uses those estimates to generate likelihood ratios. LRmix Studio was designed and developed by Hinda Haned and Jeroen de Jong, and was partly supported by a grant from the Netherlands Genomics Initiative/ Netherlands Organization for Scientific Research (NWO) within the framework of the Forensic Genomics Consortium Netherlands.

Questions regarding the software should be addressed to help@lrmixstudio.org.

2. Features

The current version of LRmix Studio has the following capabilities:

- LRmix Studio can be used to compute likelihood ratios for DNA profiles characterized with autosomal STR kits,
- LRmix Studio was thoroughly tested and validated for propositions involving at most four unknown contributors, and up to a total of five contributors,
- The hypothesized contributors under the prosecution and the defense hypotheses are assumed to be unrelated to each other, however, they can be related to an unknown contributor under the defense hypothesis,
- LRmix Studio can be used to compare any number of replicates obtained from a specific DNA sample, to any number of reference profiles. However the software has been thoroughly tested and validated for at most three reference profiles, and up to five replicates,
- LRmix Studio implements the model described in Haned et al (FSIG 2012). Optimal use of the software requires reading the relevant literature and the provided tutorial and training materials. Uncommon and/or untested scenarios may lead to unreliable results.

The current version of LRmix Studio *cannot*:

- LRmix Studio cannot be used to analyze sample profiles where the reference profiles have missing data,
- LRmix Studio cannot be used to deconvolute mixtures as it does not incorporate peak height information explicitly.

3. Tutorial

LRmix Studio is compatible with any platform having Java version ≥ 8 . The current version of the software, and future updates, are only distributed on <u>lrmixstudio.org</u>.

3.1. Import sample profiles

The first step in using the software consists in uploading the sample profiles. LRmix Studio can read files in LRmix and Genemapper® IDX formats (with or without peak heights). Any marker can be accommodated by the software, provided the same name is used in all files, although the software is not case-sensitive, it is sensitive to spaces inside maker names: if for example Penta D is used in the sample file, and PentaD is used in the reference file, these will be considered as different markers, however pentad, PenTaD, and PENTAD are considered to be the same marker.

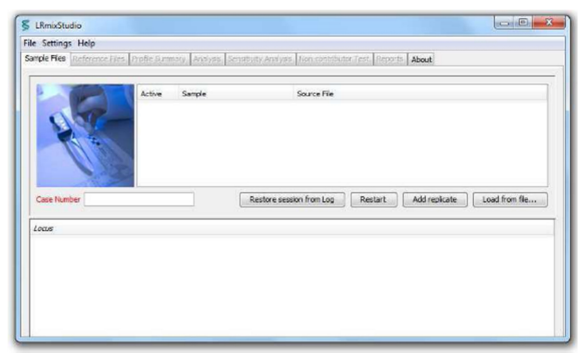


Figure 1. LRmix Studio start-up window.

Buttons

- Load from file chooses the folder from which the user wants to upload the crimesample files,
- Case Number is filled automatically from the name of the folder used to store the case files, but it can be changed to any other name by the user,
- **Restart** the software and upload a new case,
- **Restore session from Log** whenever the software is used, a log file is produced and placed in the log folder, created in the case folder. The log files are text files that contain all settings and results of an analysis. They can be uploaded to restore previous sessions so that the analysis can be redone.
- Add replicate enables adding a sample profile manually.

Once the sample is uploaded, a window displaying the alleles in the samples is obtained (Figure 2). Note that once the sample profile is uploaded the next *Reference* tab of the software that was previously grayed, becomes accessible to the user.

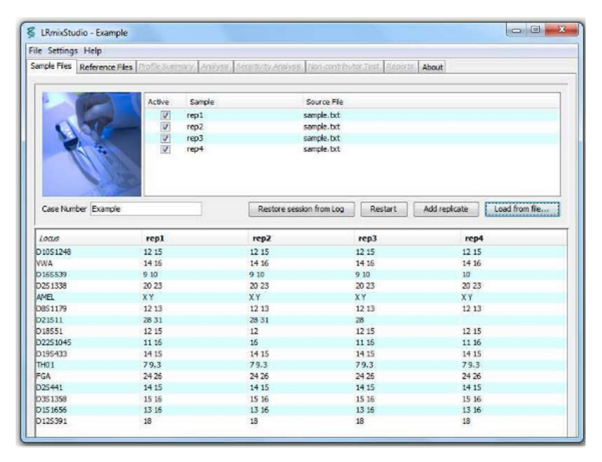


Figure 2. Upload profiles and select replicates. Once the profiles are uploaded they are displayed by the software. Note that if some replicates are not to be used in the analysis they can be un-selected at this stage.

3.2. Import or add reference profiles

Once the sample profiles are imported, the user can import the profiles of the individuals of interest (reference profiles) (Figure 3).

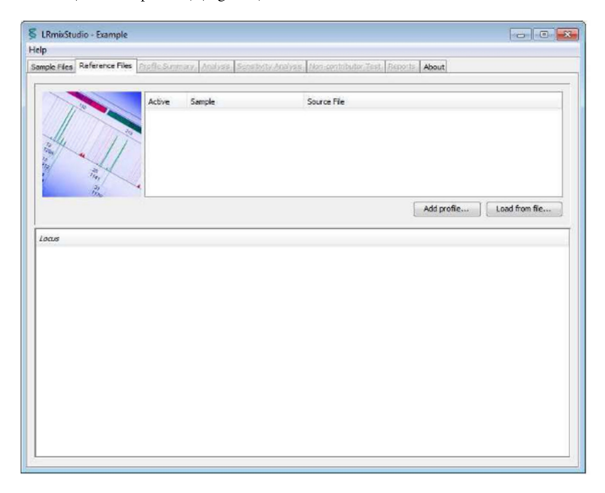


Figure 3. Upload or add reference profiles.

Buttons

- Load from file choose the folder from which you want to upload the reference-sample files, the files have to be in the LRmix format, multiple files can be uploaded at once
- Add profile in case a reference profile has to be added manually

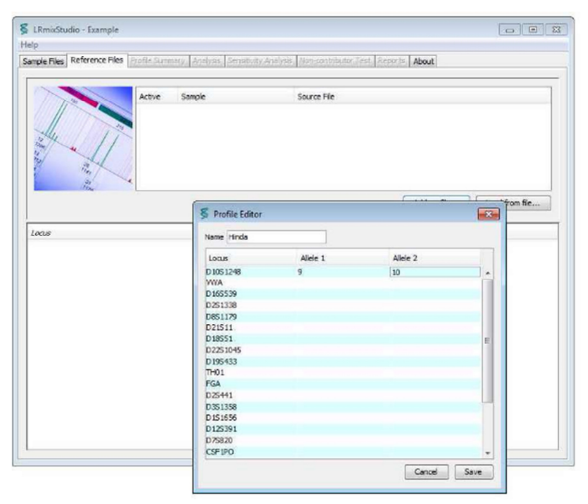


Figure 4. Reference profiles can be added manually and saved to a folder.

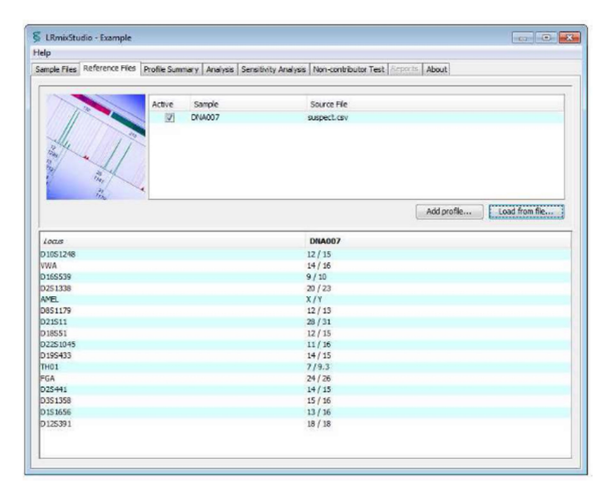


Figure 5. Reference profiles display in LRmix Studio.

When loading a profile that has a single allele at any given locus, the program will ask if this is to be converted to a homozygotic locus:

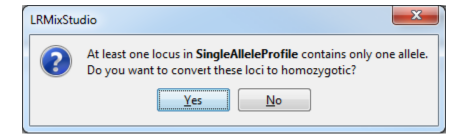


Figure 6. Dialog asking if single-allele loci should be treated as homozygotic.

Click *Yes* to convert the loci in question to homozygotic loci by duplicating the single allele present. Click *No* to leave the loci as single-allelic loci. Any duplicated alleles are highlighted in the profile view:

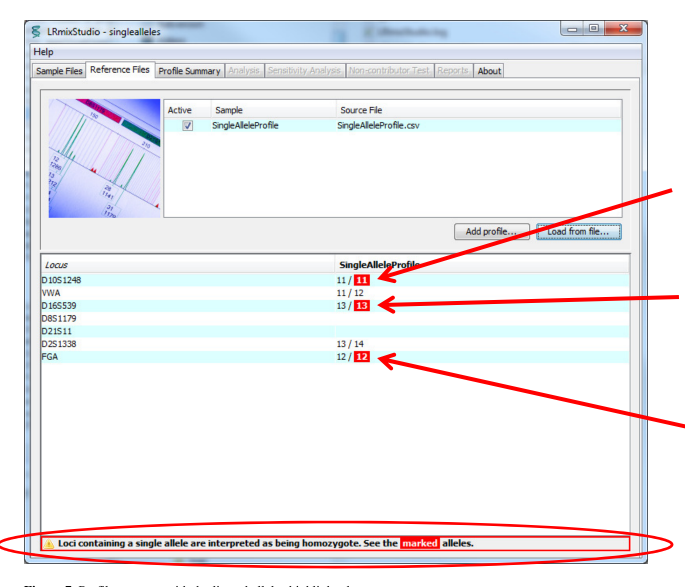


Figure 7. Profile contents with duplicated alleles highlighted.

Important note on contributors and non-contributors

Note that when the reference profiles are uploaded, only profiles relevant to the LR analysis should be uploaded, if some of the reference profiles are uploaded but later not used in the analysis tab, they will be considered as non-contributors. Non-contributor profiles can influence the likelihood ratios calculations if the Fst (θ) correction is not nil (Curran et al FSI, 2005).

3.3. Profile summary

The profile summary is an aid to the user that help visualize the alleles present in the sample profiles and those present in the reference profiles. Several filters can be used to highlight different information:

- alleles that appear in the replicates but not in the reference profiles: this filter can help highlight the alleles that might be spurious or might belong to unknown contributors,
- alleles that appear in the replicates and also in a given reference profile: this filter helps detect the allele drop-out,
- alleles that match between the reference profiles: this highlights allele sharing between the different contributors,

The different filters can be printed using the *print* button.

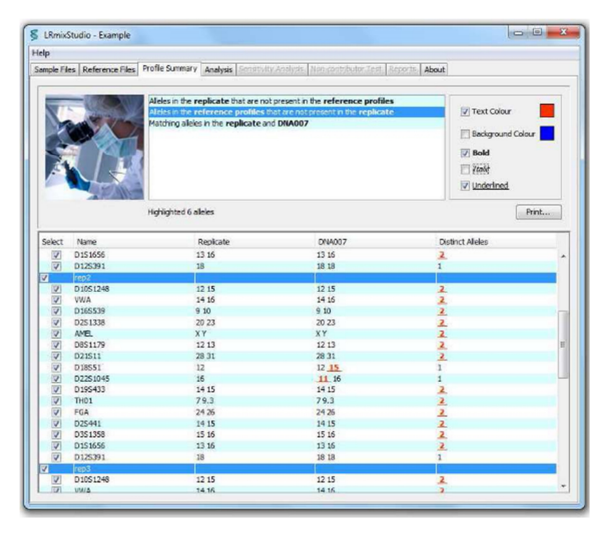


Figure 8. Profile summary.

3.4. Analysis

In the *Analysis* tab, the user can define the hypotheses of the prosecution and the defense. Under each hypothesis, the user has to define:

- the names of the contributors: the names are assigned based on the information present either within the reference files uploaded in the Reference profiles tab, or according to the name given by the user if the profiles were added manually using the editor,
- the number of unknown contributors (limit is four unknowns),
- the drop-out probabilities: for each donor, and for the unknowns,
- the Fst or theta-correction value,
- the drop-in probability (maximum is 0.50),
- the file of allele frequencies to be used,
- the rare alleles frequency to be used in case rare alleles (not in the provided file) are detected.

Analysis step 1: default screen LRmixStudio - case-01.1 Help Sample Files | Reference Files | Profile Summary | Analysis | Sensitivity Analysis Prosecution Hypothesis Defense Hypothesis Contributor Dropout Probability Contributor Dropout Probability 1 DNA007 0.1 DNA007 0.1 0 🜲 Unknown Contributors Dropout Probability for unknowns 0.1 One of the unknowns is a relative Unknown Contributors 0 💠 → of DNA007 Parent/Child w Dropout Probability for unknowns 0.1 **Parameters** Allele Frequencies Rare allele frequency 0.001 0.05 Theta correction 0.01 💠 Drop-in probability Max Threads 16 💠 Results Stop Locus Overall Likelihood Ratio Show log

Figure 9. Analysis window with the default settings.

Analysis step 2: the user defines the hypotheses and other relevant parameters

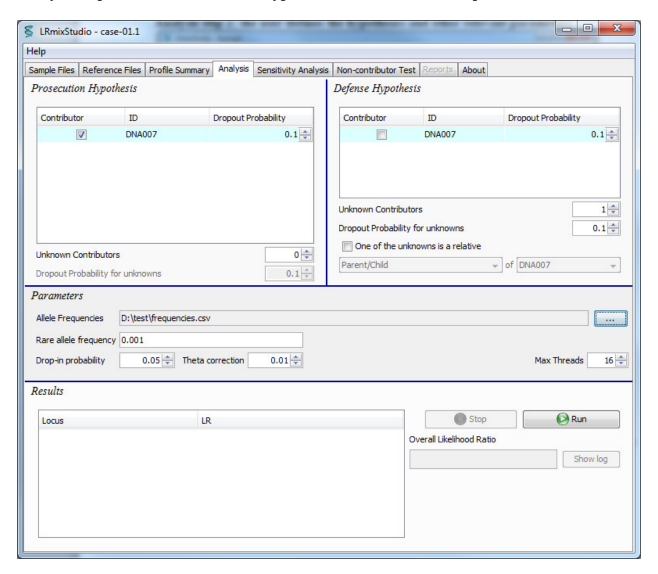


Figure 10. Analysis window where the user defined the hypotheses and the parameters. In this example, the user is evaluating the following hypotheses: Hp: Suspect (drop-out=0.10) is the donor. Hd: an unknown person (drop-out 0.10), unrelated to the suspect, is the donor. Fst is 0.01 and the drop-in probability is 0.05. The allele frequencies are that of the NGM kit in the Dutch population. If there is a rare allele in the crime-sample profile (not in the frequency file), the user chose here to assign a frequency 0.001 to it.

If one of the unknowns under the defense hypothesis is a relative of one of the profiled individuals, then the grayed box can be checked, and a relationship between one unknown under Hd and a given profiled person under Hd be selected. The formulas used for accounting for relatives follow the work of Buckleton & Triggs (see References section).

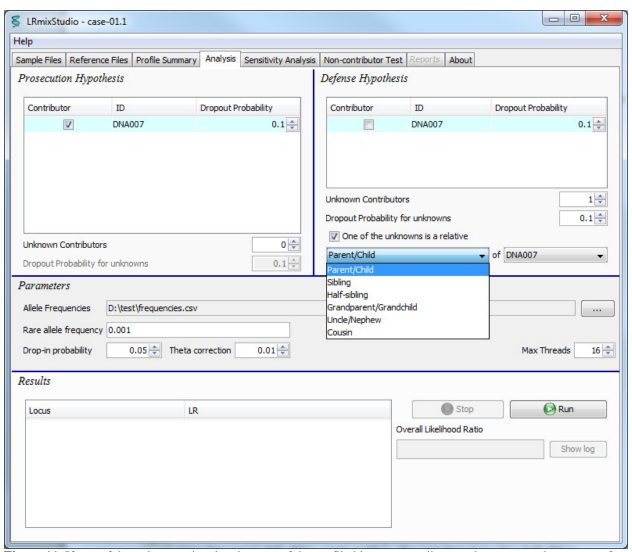


Figure 11. If one of the unknowns is related to one of the profiled known contributors, the user can choose one of several relationships: parent/child, sibling, cousin, and three other equivalent relationships: half- siblings, grandparent/grandchild, and uncle/nephew.

Analysis step 3: run the LR calculation

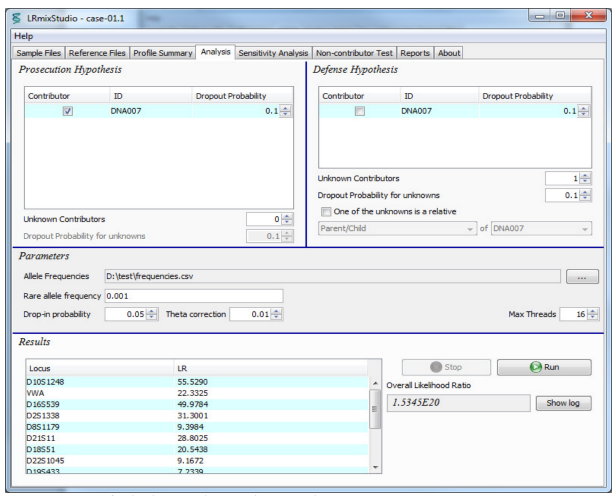


Figure 12. Result of a likelihood ratio analysis, no relatives are assumed.

If at this point the user wishes to save the results, he can go to the *Reports* tab, and save the analysis carried out so far. The report functionality is further described below.

Note that the *Show log* button has now become enabled. Whenever an analysis is performed, a log file is generated. By clicking the Show log button, the relevant logfile is displayed.

3.5. Sensitivity analysis

The sensitivity analysis (SA) plots the log10 likelihood ratios, along with the separate likelihoods of the prosecution and the defense hypotheses. The propositions evaluated in the sensitivity analysis, are those defined by the user in the Analysis tab. The drop-out parameters for the known and unknown contributors are defined in the previous step too.

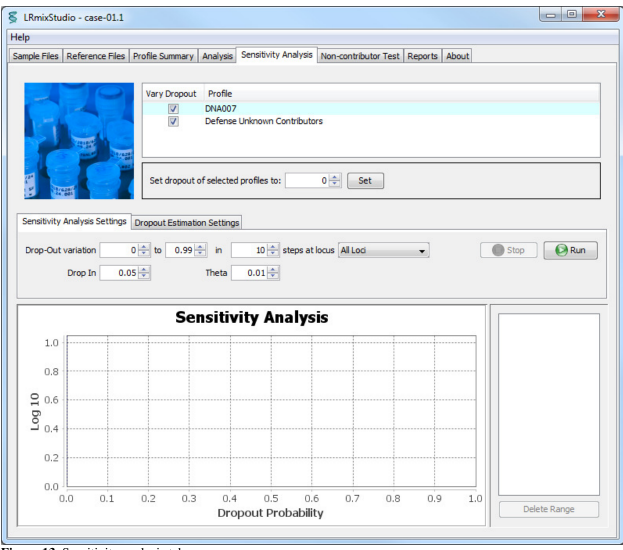


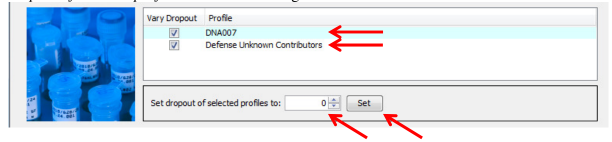
Figure 13. Sensitivity analysis tab.

Vary drop-out

The user can first choose which of the known and unknown contributors will have a drop-out probability that is varied in the SA. In the example below, both *DNA007* and *Defense Unknown contributors* are checked. This means that all contributors will have the same dropout probabilities in the SA. If the drop-out of the major should not be varied in the analysis, as it is the case in this example, the box should be unchecked.

Setting dropout of selected profiles

Once a value for dropout has been arrived at through the means of Dropout Estimation and Sensitivity Analysis, this value can be set directly to the selected profiles by entering it in the *Set dropout of selected profiles* field and clicking the *Set* button:



It is of course also possible to switch back to the Analysis tab and set the dropout for relevant profiles manually.

Sensitivity Analysis Settings

The default variation ranges are zero to 0.99, and the user can choose at most 100 values. The SA can also be performed for a given locus, while others would be ignored (All loci button). This can help understanding the relative contributions of different loci. the drop-in and the Theta-correction values are set at the values chosen in the *Analysis* step, however the user can change them during the SA. The *Run* button runs the SA.

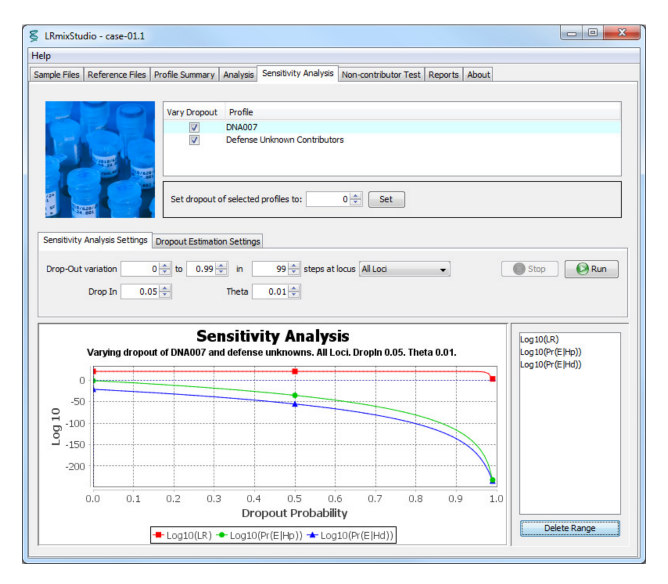


Figure 14. Result of a sensitivity analysis for the Example case. Note that right-clicking on the plot frame allows to zoom in and out.

The output of the SA is displayed in Figure 14 above. If a new analysis is carried out, with a different parameter, for example, a different drop-in rate, then the curves are displayed on the same graph. The right panel allows selecting the relevant curves for more clarity (delete range button).

The results of the sensitivity analysis are stored in a log-file, and they can also be printed into the report.

If the evidence does not support the configured hypotheses, the results for all dropout values may be non-numerical (Infinity etc.). In this case the following error will be displayed:

"The Sensitivity Analysis did not result in any numerical results. All LRs were either 0, Infinity or not representable as a number. Please check your hypotheses."

Drop-out estimation Settings

This tab allows the user to estimate the drop-out probability following the method described in H. Haned et al (2012). This is a qualitative estimator of the drop-out probability of the whole profile, based on the average numbers of alleles observed in the profile. The user can choose the number of drop-out values to explore between 0 and 0.99, as well as the level of drop-in. If there are fixed individuals that have no drop-out, then the check box in the top frame has to be unchecked accordingly. The output is an interval of the plausible range of drop-out, plotted on the SA, but also displayed as a highlighted area in the plot.

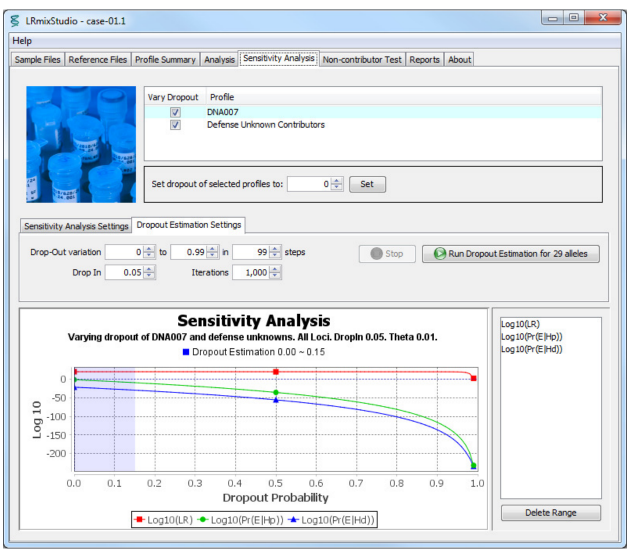


Figure 15. Result of a sensitivity analysis and drop-out estimation for the Example case.

In case the tested scenario assumes a total number of contributors that is not supported by the qualitative estimator of the drop-out, then the following error is obtained: "drop-out estimation resulted in no matching attempts under prosecution". This might happen if the average number of alleles across the replicates, say 50, is not supported by the hypothesis of a single-source sample.

3.6. Non-contributor tests

Non-contributor tests are an optional aide, meant at assisting the understanding of the case-specific likelihood ratio (Gill & Haned, 2013). The tests consist in calculating the LR for the propositions chosen by the user, where the profile of the person of interest is replaced by a random profile generated from sampling alleles with respect to their frequencies outlined in the file of allele frequencies (provided in the *Analysis* tab).

Given the parameters and hypotheses chosen in the *Analysis* tab, the non-contributor tests consist in calculating the LR obtained when replacing the profile of the person of interest, by the profile of a simulated random man.

This is carried out n times, where n is the number of *iterations* defined by the user. The output of the test is a distribution of n (log10) likelihood ratios, which are represented in a barplot as follows:

- the case-specific log10 LR, obtained with the person of interest, is displayed in red,
- the minimum, the maximum, the 1%, the 50% and the 99% percentiles of the obtained distributions are displayed in grey.

Figure 16 below gives an example of the non-contributor tests carried out for case 10.

Important notes

- 1. For the non-contributor tests to yield a result, the drop-out and drop-in probabilities (defined in the *Analysis* tab) must be different from zero, see details in Gill & Haned (2013),
- 2. The randomly generated profiles are created using random sampling of the alleles and their frequencies, as given in the population frequencies file. These profiles are generated at random, and no relatedness is assumed, even if the Hd hypothesis involves relatives.

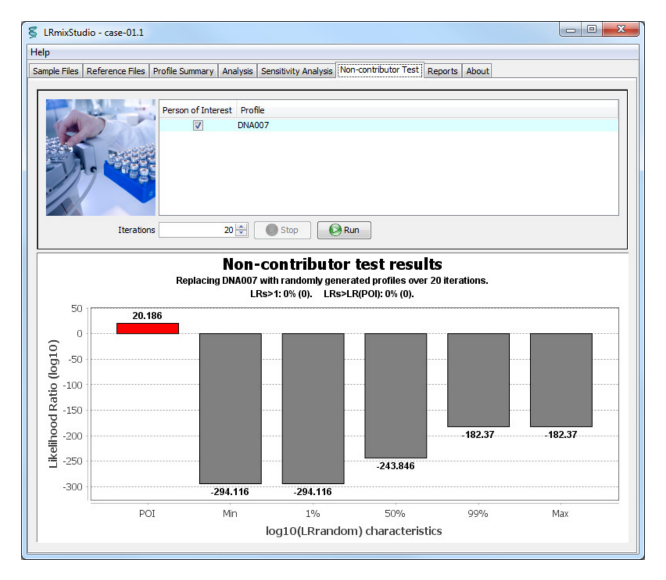


Figure 16. Result of the non-contributor tests for the Example case, where the suspect is replaced by the randomly generated profiles. The details of the non-contributor tests are printed to a log file stored in the folder of the case being analyzed.

3.7. Automatic report generation

The *Reports* tab describes all the analyses carried out by the user, within a given session. The user can select the analysis to be exported in a report, in a PDF format. Multiple analyses can be exported to the same report. Note that, in addition to the report, log files are automatically generated and stored in a *log* folder, within the case folder that contains the case files. The log files contain all the actions and results obtained by the user within a given session.

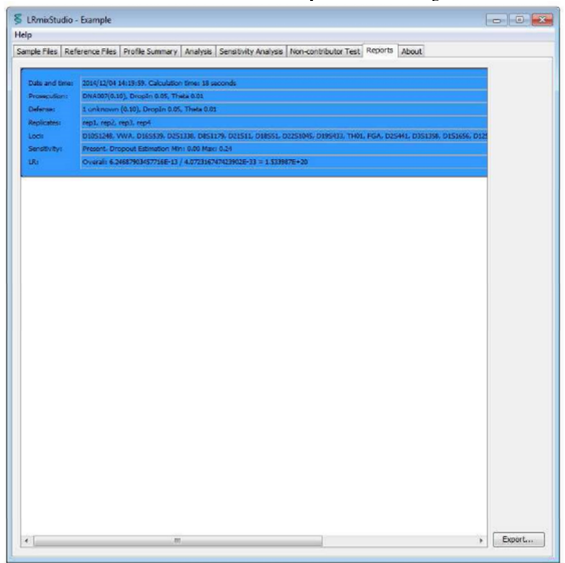


Figure 17. Select the analysis to be exported to a the PDF report.

Once the **Export** button is pressed, a comment window pops-up (Figure 18), where the user can add comments to the report. Comments are optional.

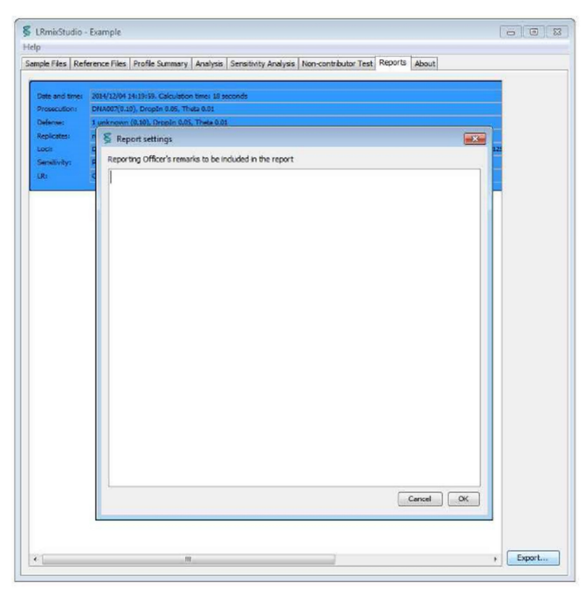


Figure 18. Report generation and the comment section.

4. How to report bugs

A bug is defined as an error, or a failure of the software that causes it to produce an incorrect or unexpected result. If such error is encountered, the following procedure has to be followed:

- prepare the log files (that are generated in your case folder, see section 3) and send them by email to help@lrmixstudio.org
- it is important that the error is described thoroughly in your email, so that the problem can be fixed quickly. Simply describing a problem will not be enough to get the help you need.

5. Join the LRmix Studio user community

Visit lrmixstudio.org/user-group, to ask questions, discuss cases or make suggestions to the development team.

References

- P. Gill and H. Haned. A new methodological framework to interpret complex DNA profiles using likelihood ratios. Forensic Sci. Int. Genet., 7(2):251-263, 2013.
- P. Gill, et al. Interpretation of complex DNA profiles using empirical models and a method to measure their robustness. Forensic Sci. Int. Genet., 2: 91-103, 2008.Z
- H. Haned et al. Exploratory data analysis for the interpretation of low template DNA mixtures. Forensic Sci. Int. Genet., 6(0): 762-774, 2012.
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