RGeasy Instruction Manual

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1 Introduction

This manual is a guide for researchers seeking validated reference genes for gene expression analysis via RT-qPCR, as well as for those aiming to register species (animals, plants or microorganisms) used in their reference gene validations studies (Figure 1). RGeasy makes research access and development simpler, as it provides greater data dissemination, reduces cost, and decreases the time required to conduct each study. In addition, due to the increased visibility of the registered studies on RGeasy database, these studies can potentially be cited more than usual, as detailed along this manual.

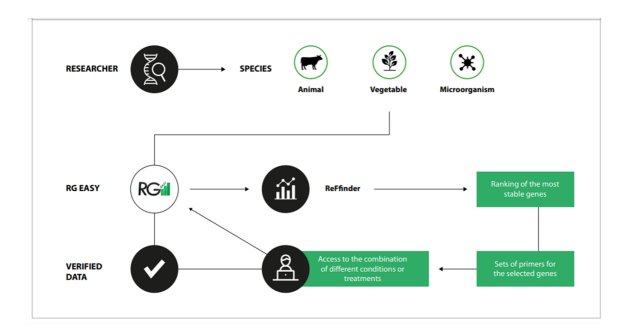


Figure 1- Maintenance and workflow of the RGeasy tool. From the registration of animal, plant or microorganism species, researchers deposit their data (Cq's values) on RGeasy database, which are immediately verified. Then, users can run all possible combinations of conditions/treatments for each study. The new combinations of treatments is ranked by RefFinder (XIE et al., 2012), and RGeasy provides, in addition to the ranking with reference genes, a set of validated primers for each reference gene.

2 Access to RGeasy database

When using RGeasy, user has access to the species registered on the tool by clicking on "Species" in the navigation bar located in the upper part of the initial interface (Figure 2).

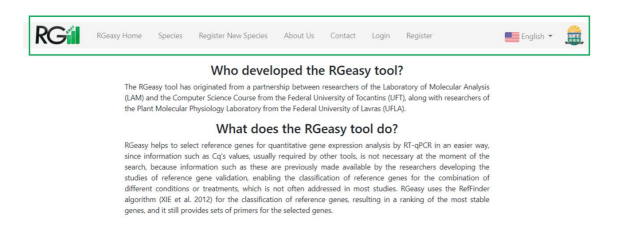


Figure 2- RGeasy's initial graphic interface

Species on RGeasy are separated into three categories: Animals, Plants, and Microorganisms (Figure 3). By clicking on the species of interest, it is automatically shown all the reference gene validation studies registered on RGeasy for that species. In this interface, the user has access to each study by clicking on its title, and under the title from each study, it is displayed the types of samples analyzed on them.

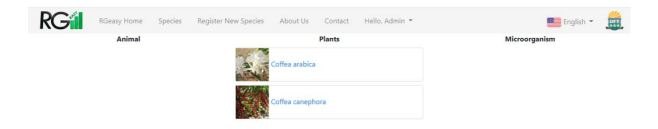


Figure 3- Species categories on RGeasy.

In order to define the desired combination of treatments or conditions, users must select the samples of interest by clicking on the icon beside them (Figure 4). The result is instantly shown by clicking on "Run RefFinder".

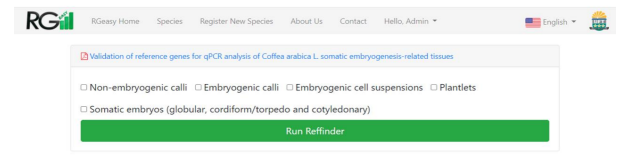


Figure 4- Samples analyzed in the study entitled "Validation of reference genes for qPCR analysis of Coffea arabica L. somatic embryogenesis-related tissues" by Freitas et al. (2017).

Since RGeasy uses the RefFinder tool to analyze the stability of the reference genes, a table is generated on the results page with the ranking of genes according to the following algorithms: RefFinder, Delta CT, Bestkeeper, Normfinder and Genorm (Figure 5).

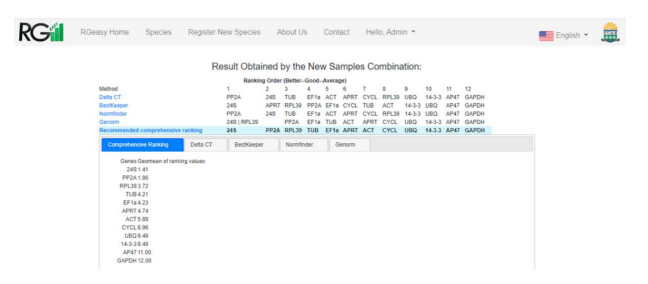
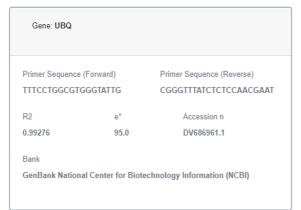


Figure 5- RGeasy's result for the different samples analyzed by Freitas et al (2017).

In addition, on the results page, RGeasy provides a table with some additional information for each reference gene, according to the stability ranking from RefFinder. For each reference gene, the primer pair, the correlation coefficient (R²), the amplification efficiency, the accession number, and the database from which the sequence was obtained, are made available to the user (Figure 6).



Primer Sequence	e (Forward)	Primer Sequence (Reverse)
TTTCCTGGCGT	GGGTATTG	CGGGTTTATCTCTCCAACGAAT
R2	e*	Accession n
0.9923	92.0	GT648763.1
Bank		
GenBank Nation	nal Center for Biote	echnology Information (NCBI)

Figure 6- Representation of the general information made available by RGeasy for two reference genes, *Ubiquitin* (*UBQ*) and *photosystem I P700 chlorophyll a apoprotein A2* (*PSAB*), present on RGeasy's database.

When selecting reference genes through RGeasy, it is essential to include, in the Materials and Methods section of the user's article, the correct form of RGeasy's citation, which is unique, according to the research being developed (Figure 7).

How to Cite

The RGeasy tool (citation soon) was used for the selection of reference genes through a new combination of treatments, obtained from the study developed by Freitas, et al. (2017) and ranked by the RefFinder (XIE et al., 2012) tool.

References:

- Reference Genes Easy
- Reffinde
- Validation of reference genes for qPCR analysis of Coffea arabica L. somatic embryogenesis-related tissues

Figure 7- The correct way of citing RGeasy in each study is present at the end of the result's page.

3 Registering new species

3.1 Registering a single species

In order to register new species, the user should click on the "Register New Species" icon located in the navigation bar (Figure 8). The user should first insert a image file that best characterizes the species that is being registered. Then, the user should add the scientific name of the species, as well as the title of the article, the DOI (Digital Object Identifier), publication year, authors names and the Cq's values of the study (Figure 8).

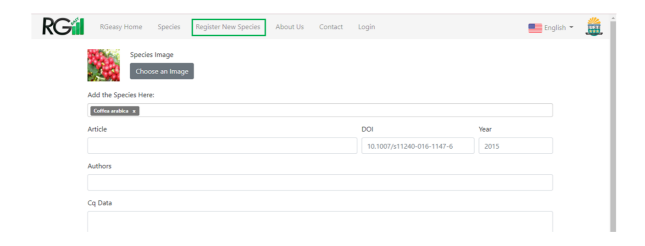


Figure 8- Registration a new species on RGeasy database.

Once each required information is filled out, the user should not press the "Enter" key, since the tool will understand that all fields have been completed and automatically generate the table displaying the Cq value (Figure 11) of the samples for each reference gene being analyzed. However, since not every field is filled out, this action will result in an error message, as shown in figure 9.

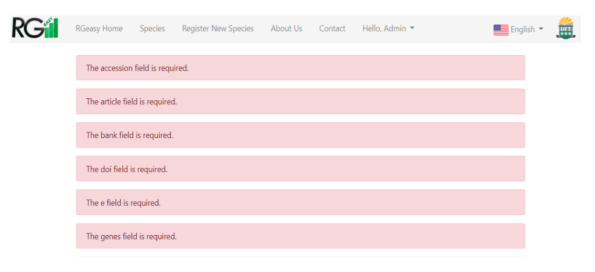


Figure 9- Error message generated due to blank fields during registration of a new species on RGeasy's database.

Author's names should be inserted in the "Authors" field and must be separated exclusively by commas (Figure 10).

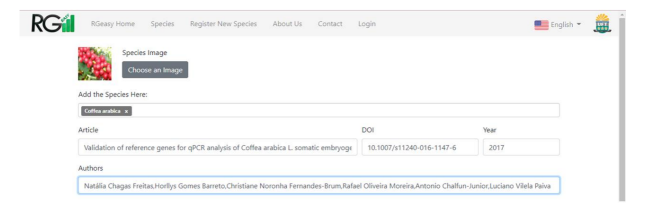


Figure 10- Filling out the author's field during registration of a new species on RGeasy's database.

After filling out the required fields (Figure 11) and pressing the "Enter" key, a table that allows the user to edit the data if necessary will be generated.

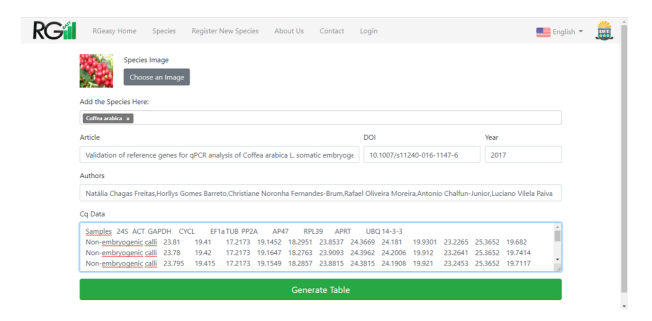


Figure 11- Registration a new species on RGeasy's database.

Once the user press on the "Generate table" icon (Figura 11), the table cannot be further edited and a pop-up window will be exhibited, asking the user to select the species category (Animal, Plant or Microorganism) of the new species that is being added (Figure 12).

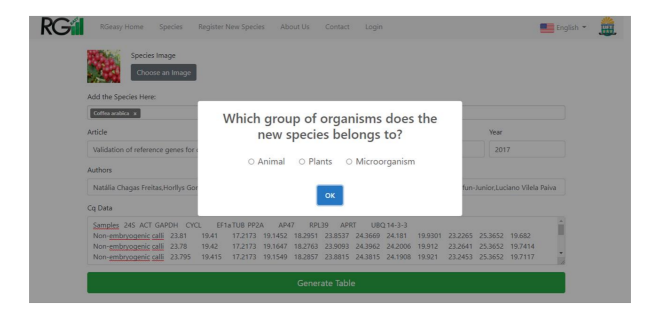


Figure 12- RGeasy's users must inform the species group of the new species being added after clicking on "Generate table".

Below the table containing the Cq values, a description from each candidate reference gene being analyzed should be given by the user (Figure 13). This information must be carefully completed, since it will be available for other researchers using RGeasy.

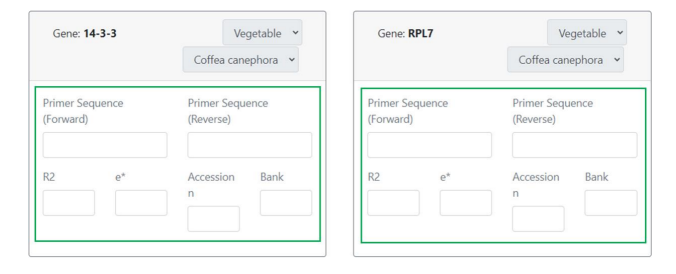


Figure 13- Gene information users must complete for each candidate reference gene of the new species being registered on RGeasy's database.

The process of registering a new species is completed when users press "Generate", once every field related to the candidate reference genes is filled out.

3.2 Registering two or more species

Researchers carrying out studies that analyze the stability of candidate reference genes for two or more species can make the registration of the different species through a single registration process (Figure 14). However, this is true only if the same group of candidate reference genes is being analyzed for each species. Otherwise, species should be separately registered and thus the article will have to be registered more than once.

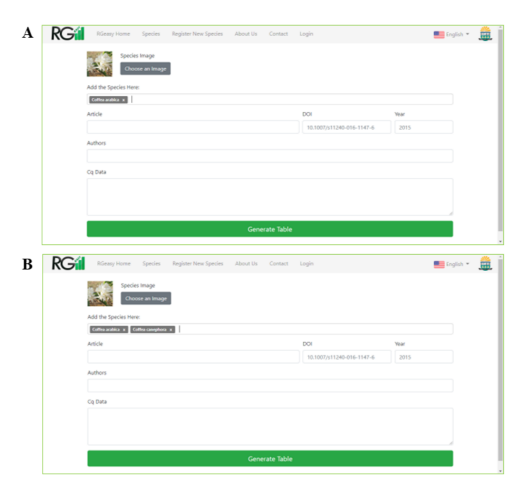


Figure 14- Registration process of two or more species. The name of the second species should be added right after the name of the first species, with these names being separated by a comma. Registration of one species (A). Registration of two or more species (B).

During the registration of the Cq values, it is essential to identify each sample with the name of the species to which it refers (Figure 15), so thus RGeasy can correctly differentiate each tissue. Otherwise, the tool may recognize two or more samples as only one sample. The further steps of registering two or more species are similar the those of registering a single species.



Figure 15- Description of how to correctly identify each sample on RGeasy. In this case, root tissue was used for two different species (*C. arabica* and *C. canephora*) (Fernandes-Brum et.all, 2017) and as highlighted in green, the identification of each sample in this case was made by writing name of the species followed by the name of the tissue.