

QuickGuide: RealFast™ Variant Detection on MIC qPCR Cycler

Setup for Variant Detection and Onco RealFast™ Assays:

- Open the micPCR software.

Definition of a new assay:

- Define a new assay by **New > Assay** in the menu bar.
- Within the **Assay Setup** choose **Information**.
 - Set the **Chemistry Type** to **Hydrolysis Probes**.
 - Name your target (e.g. the name of the gene/allele of interest) and optionally provide a Description.
 - Define the **5´** and **3´ Modifier** in the field for **Probes**. Typically, the gene/allele of interest is labeled with 5´-FAM and 3´-BHQ1.
 - Klick on “+” next to **Targets** and define the second probe, which targets the internal PCR control. Typically, the PCR control is labeled with 5´-HEX and 3´-BHQ1.

Chemistry Type

+ Targets

Name	<input type="text" value="HLA-B27"/>		
Description	<input type="text"/>		
	Amplicon Length <input type="text" value="0"/>		Sequence <input type="text"/>
Forward Primer	Name <input type="text"/>		Sequence <input type="text"/>
Reverse Primer	<input type="text"/>		
Contains Alleles	<input type="checkbox"/>		
Probes	Name	5´ Modifier	3´ Modifier
	<input type="text" value="New Probe"/>	<input type="text" value="FAM™"/>	<input type="text" value="BHQ® - 1"/>

Name	<input type="text" value="Control"/>		
Description	<input type="text"/>		
	Amplicon Length <input type="text" value="0"/>		Sequence <input type="text"/>
Forward Primer	Name <input type="text"/>		Sequence <input type="text"/>
Reverse Primer	<input type="text"/>		
Contains Alleles	<input type="checkbox"/>		
Probes	Name	5´ Modifier	3´ Modifier
	<input type="text" value="New Probe"/>	<input type="text" value="HEX™"/>	<input type="text" value="BHQ® - 1"/>

» **Note:** For assignment of fluorophores to hydrolysis probes see *Instructions for Use of the respective RealFast™ Assay*. «

- Within the **Assay Setup** choose **Profile**.
 - Define in **Temperature Control** the mode **Fast TAQ (v3)** and in **Volume** 20µl.
 - Setup the PCR program according to the *Instructions for Use of your RealFast™ Assay*.
Data acquisition should be by default at the end of the 60°C step (camera symbol is black).

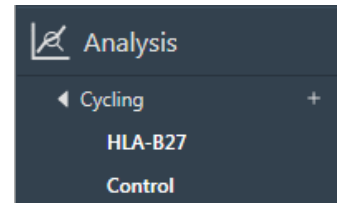
- **Safe** your assay!

Setup a new run:

- Define a new run by **New > Run** in the menu bar.
 - In the menu on the left side choose **Assays** and click on “+” next to it.
 - Via the **Shortcut Library** you can choose one of your previously designed assays for variant detection.
- Within **Run Setup** choose **Samples** and name your samples.
 - In the field **Type** choose **NTC** for your No Template Control (NTC), **Positive Control** for your positive control, **Negative Control** for your negative control, and **Unknown** for the samples you wish to analyze.
 - In the field **Assay** choose the correct variant detection assay
- Save the run setup, load your PCR tubes and start the run.

Analysis of Variant Detection and Onco RealFast™ Assays:

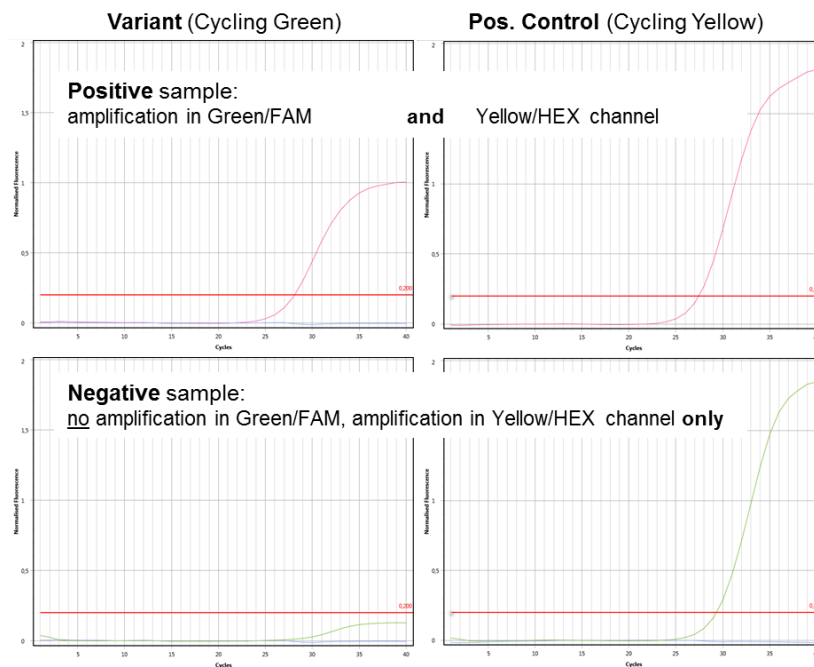
- Open the micPCR software.
- Click on the **Open** icon and select the data file you wish to analyse.
- Add a new **Cycling** analysis by clicking “+” in the **Analysis** section. In the drop down menu you can select the targets, which you previously designed.



- Select the target corresponding to the internal **PCR control** in the menu on the left.
- Set **Ignore first cycles** to “10” within **Parameters** (optional). Choose **Dynamic** as **Method**. The **Exclusion** method should be **None**. Set the threshold in the exponential phase of the amplification curves.



- Verify that all samples and the **Positive Control** show an amplification curve in the Yellow/HEX channel. Mouseover a sample in the **Samples** section highlights the corresponding amplification curve.
- Select the target corresponding to your **gene of interest** in the menu on the left.
- Set **Ignore first cycles** to “10” within **Parameters** (optional). Choose **Dynamic** as **Method**. The **Exclusion** method should be **None**. Set the threshold in the exponential phase of the amplification curves.
- In the **Cycling Analysis** window, only samples which are positive for the variant will show an amplification curve in the Green/FAM channel. Move the mouse over a curve to see the sample name.



» **Note:** For **Onco RealFast™** Assays (e.g. EGFR T790M,...), it is **mandatory** to do the analysis according to the section *Data Analysis / Interpretation of Results* in the respective *Instructions for Use*. «

- Optionally generate a report by clicking “+” next to **Reports** on the left.