

# **FLIM FRET Calculation for Multi Exponential Donors**

## Summary

This tutorial shows step-by-step, how the "Lifetime FRET Image" analysis of SymPhoTime 64 can be used to calculate pixel-by-pixel the average FRET efficiency in an image containing several cells transfected with Cerulean/YFP-constructs. To calculate the average FRET-efficiency, first the donor only lifetime is determined using cells that have only been transfected with the donor construct by performing a ROI-fit with the FLIM-analysis. The amplitude weighted average donor lifetime is used to calculate the average FRET-efficiency using the "Lifetime FRET Image" analysis.

## **Background information**

FLIM-FRET is a robust method to determine the FRET efficiency of a suited donor acceptor pair. If the FRETdonor molecule has a more exponential decay kinetic, the amplitude weighted average lifetime has to be used to calculate the average FRET efficiency. The script usage is demonstrated on an image of several cells which have been transfected with CENP-B-Cerulean and YFP-CENP-A. Cerulean is a CFP variant an frequently used as a FRET donor. The CENP-complex consists on several proteins and is involved in mitosis. For further details on the sample, see S. Orthaus et al., Chembiochem. 2008, 9, 77-92, and the PicoQuant application note http://www.picoquant.com/images/uploads/page/ files/7267/appnote\_flim\_fret.pdf. The example image used is an image showing four cells, two of them only transfected with the CENP-B-Cerulean complex, the other two transfected with both constructs. Therefore, the same image can be used to determine the donor only lifetime and also to perform the pixel-by-pixel fit to calculate the FRET efficiency.

## **Step-by-Step Tutorial**

## Determine the donor only lifetime using the FLIM script

- Start SymPhoTime 64 software.
- Open the "Samples" workspace via "File\open Workspace" from the main menu.

## Note:

The "Samples" workspace is delivered with the SymPhoTime 64 and on the CD-ROM-drive and contains example data to show the function of the SymPhoTime data analysis. If you haven't installed it on your computer, copy it from the the DVD onto a local drive before going through this tutorial.

## **Response:**

The files of the sample workspace are displayed in the workspace panel on the left side of the main window.



• Highlight the file CENP-labelled \_cells\_for\_FRET.ptu by a single mouse click.



• Select the "Analysis" tab and in there, open the drop down menu "Imaging".



#### Note:

The drop down menu can be opened and closed by clicking on the grey button on the left side of the header of the drop down menu:

• Start the FLIM script by clicking on "Start".



#### **Response:**

The FLIM script is applied to the file CENP-labelled\_cells\_for\_FRET.ptu. Thereby, a new Window opens:



#### Note:

The window contains five different regions:

**Upper left:** Imaging analysis options.

**Upper center:** Fast FLIM image, displayed in false color scale. The brightness encodes the intensity while the color encodes the average "Fast FLIM" lifetime, i.e. the mean arrival times of the photons after the laser pulse. When not defined otherwise, intensity and color scale stretch from minimum to maximum. As the mean arrival time of the background photons in the areas where no fluorescence is detected, is randomly spread over the TCSPC window, the mean photon arrival time of the dark background is usually very long (up to 1/2 of the TCSPC window), which makes the color scale loaded by default sometimes unsuited for displaying the lifetime contrast in the actual sample. In this case, the scale has to be adapted.

**Upper right:** Fluorescence Lifetime Histogram. Here the frequency of photon counts corresponding to the individual mean lifetimes is plotted. The settings of the plot can be adapted using the controls on the right of the plot.

**Lower left:** Lifetime fitting parameters. Here lifetime fitting models can be defined. Fitting is then applied to the lifetime graph(s) in the TCSPC histogram.

**Lower center/left:** By default, this set shows the TCSPC histogram of all photons in the image in green (= dataset 0) and the TCSPC histogram of a single pixel in grey (= dataset 1). In red an estimation of the instrument response function (=IRF) is shown. The IRF reconstruction is deducted from the rising edge of the TCSPC histogram.

• As data channels, check only Channel 2, as in this channel the donor emission has been recorded.

FLIM			
Binning:		1 Points	<b>A</b>
Set Time Gate:			
			~ 
Select Data Channels: 1:	2:	3:	4:
Threshold		50 Cnts	
Calculate FastFLIM		FLIM Fit	

• Press "Calculate Fast FLIM" to update the image.

## **Response:**

The image and the graphs are recalculated, using photons only from channel 2.



• To enhance the lifetime and intensity contrast, set in the image the intensity from 1 count to 30 counts and the lifetime from 0 to 5 ns.

### **Response:**

The image scales are adapted.

### Note:

Now four cells are visible, the two nuclei on the left have longer Fast lifetimes then the other two. These are the cells transfected only with the donor constructs.

CENP-labelled_cells_for_FRET.ptu Channels: 2	🔵 Gray 💿 Rainbow 💿 RGB
	Events[Cnts]
	Min 1 Max 30 A
the second se	
	> >
100 µm	> >
· · · · · · · · · · · · · · · · · · ·	

• Keep the mouse pointer over the image and open the context menu by a right mouse click. Select the "Free ROI" tool.



## Note:

For more information about the different ROI selection tools, check the tutorial ROI fitting using the FLIM script.

• Draw a ROI around one of the left cells by maintaining the left mouse button activated while circling the selected part of the image. Then release the left mouse button, press "Shift" continuously and cycle the other left nucleus. This way, the donor only cells are marked as ROI 0.

## **Response:**

• Only the selected area is now shown in color.



• In the TCSPC window, the photons from ROI 0 are added in grey.



• In the decay form on the lower left, select n-exponential reconvolution as fitting model.

Fitting Model:	n-Exponential Tailfit Help	
Decay:	n-Exponential Tailfit n-Exponential Reconvolution	
IRF: Import	Calculated IRF	Remove
Model Parameters:	n 1 📥	

#### **Response:**

- In the TCSPC window, the IRF is displayed in bright red, the data fitting limits are moved to the border of the TCSPC window.
- The new fitting parameters "Shift IRF" and "Bkgr IRF" (=background IRF) appear in the fitting parameter

#### table.



#### Note:

The software offers the possibility to fit the data using a n-exponential tailfit or a n-exponential reconvolution fit. A tailfit can be used when the fitted lifetimes are significantly longer than the instrument response function. Still a reconvolution fit is usually preferable, especially for FRET determinations, because the complete decay is fitted, while for a tailfit, the start of the fitting range is usually a bit arbitrary, and the short component can be underestimated. For explanation on the fitting model and the used equations, click on the "Help" button next to the selected model. This opens a help window containing the fitting equation and the explanation of the different parameters.

• Under "Decay" select "ROI 0".

#### **Response:**

The TCSPC curve of ROI 0 is highlighted in green.



• At the IRF, click on the "Import" button.

Fitting Model:	n-Exponential Reconvolution   Help
Decay:	ROI 0
IRF: Import	Calculated IRF Remove
Model Parameters	n 1 📥

#### **Response:**

A window opens, where the file containing the IRF can be selected.

• Select the file CENP-labelled\_cells\_for\_FRET\_IRF\_Det2.ptu and press "OK".

## Note:

This file contains a measured instrument response function (IRF) of detector 2. The best way to measure an instrument response function is measuring a dye that is highly quenched in a saturated solution of KI. See e.g. Szabelski et al., Appl. Spectrosc. 2009, 63, 363-368.



## **Response:**

The decay of the imported IRF is shown in grey in the TCSPC window.

• Activate the imported IRF by selecting the IRF file in the IRF drop down menu.

Fitting I	lodel:	n-Exponential Reconvolution	Help
Decay:		ROI 0	
IRF:	Import	Calculated IRF	Remove
Model F	Parameters:	Calculated IRF CENP-labelled_cells_for_FRET_IRF_Det2.p	

#### **Response:**

The imported IRF is highlighted in red.



• As model parameters, select n="2".

## Note:

In this case, the decay is double exponential. As it is not the scope of the tutorial to explain standard lifetime fitting (see the tutorial "ROI fitting using the FLIM script"), we skip the steps of finding the correct fitting model here.

Fitting Model:	n-Exponential Reconvolution	
Decay:	ROI 0	
IRF: Import	CENP-labelled_cells_for_FRET_IRF_D	emove
Model Parameters:	n 2 🔺	
Parameter	Value	Fit 🔺
A [1] [kCnts] Limit	s 0,000 ± 0,000	<b>₹</b> √_
A [2] [kCnts] Limit	s 0,000 ± 0,000	ŧ٠
τ[1] [ns] Limit	s 0,000 ± 0,000	<b>₹</b> <
τ[2] [ns] Limit	s 0,000 ± 0,000	<b>₹</b> <
Bkgr Dec [Cnts] Limit	s 0,000 ± 0,000	<b>₹</b> <
Shift <sub>IRF</sub> [ns] Limit	s 0,000 ± 0,000	<b>€</b> <
Bkgr IRF [Cnts] Limit	s 0,000 ± 0,000	<b>₹</b> <
I[1] [kCnts]	0,000 ± 0,000	
I[2] [kCnts]	0,000 ± 0,000	
Isum[kCnts]	0,000 ± 0,000	
A sum [kCnts]	0,000 ± 0,000	
τ Av Int [NS]	0,000 ± 0,000	
τ Av Amp [NS]	0,000 ± 0,000	

• Click on "Initial Fit".

**Response:**The fit is calculated.



• The software calculates the amplitude weighted average lifetime  $\tau_{Av.Amp.}$  of 2.675. This is the result of the average donor lifetime in absence of FRET.

#### Note:

The fitted lifetime can slightly differ depending on the ROI drawn. It is important to use the amplitude weighted average lifetime. While the intensity weighted average lifetime corresponds to the real average arrival time of the photons and is therefore also close to the Fast lifetime in case of low background noise, the amplitude weighted average lifetime reflects the quenching of the donor due to the FRET process and therefore the total drop of intensity. To see the definitions of both averages, click on the "Help" button next to the fitting model.

• Save the result by clicking on "Save Result".

Analysis		
Region of Interest		
Frame		
FLIM		
Binning:	1 Points	▲ ▼
Set Time Gate:		
		¥.
Select Data Channels: 1:	2: 🗹 3: 🗌 4:	
Threshold	50 Cnts	
Calculate FastFLIM	FLIM Fit	
▲ File		
	Save Result	
Save Defaults	Restore Defaults	

#### **Response:**

A result file (FLIM.pqres) is stored under the raw data file (CENP-labelled\_cells\_for\_FRET.ptu).



• Now the first part is finished and the FLIM window should be closed.

## Calculate the FRET efficiency image using the FLIM-FRET script

• Highlight the file CENP-labelled \_cells\_for\_FRET.ptu again by a single mouse click.



• Select the "Analysis" tab and in there, open the drop down menu "Imaging".



• Start the "Lifetime FRET Image" script by clicking on "Start".

Lifetime FRET Image	
A lifetime based FRET image. Select an image TTTR file and press 'Start'.	
Help Start	

#### **Response:**

The "Lifetime FRET Image" script is applied to the file CENP-labelled \_cells\_for\_FRET.ptu. Thereby, a new Window opens:



The window contains five different regions: Upper left: Imaging analysis options

**Upper center:** Fast FLIM image, displayed in false color scale. The brightness encodes the intensity while the color encodes the average "Fast FLIM" lifetime, i.e. the mean arrival times of the photons after the laser pulse. When not defined otherwise, intensity and color scale stretch from minimum to maximum. As the mean arrival time of the background photons in the areas where no fluorescence is detected, is randomly spread over the TCSPC window, the mean photon arrival time of the dark background is usually very long (up to 1/2 of the TCSPC window), which makes the color scale loaded by default sometimes unsuited for displaying the lifetime contrast in the actual sample. In this case, the scale has to be adapted.

**Upper right:** Fluorescence FRET Efficiency Histogram. As the FRET efficiency image has not been calculated yet, this plot is still empty. Later, the settings of the plot can be adapted using the controls on the right of the plot.

**Lower left:** Binding histogram. As the FRET efficiency image has not been calculated yet, this histogram is also still empty. As this diagram can only be calculated if the FRET donor dye has single exponential decay kinetics, changing the fitting model results in an exchange of this graph to a FRET distance histogram.

**Lower center/left:** TCSPC histogram of all photons in the image in green (= dataset 0) and the TCSPC

histogram of a single pixel in grey (= dataset 1). In red an estimation of the instrument response function (=IRF) is shown. The IRF reconstruction is deducted from the rising edge of the images TCSPC histogram. On the left of this graph, the fitting model and parameter controls are displayed.

- Select only channel 2 as active channel.
- As the photon number/pixel is quite low in this image, select a Pixel Binning of 4 points.
- Press "Calculate FastFLIM" to update the image.



#### **Response:**

The pixel binning is applied to the image and the image is recalculated, using only photons from detector

## channel 2.

	FLIM Image	
	Events[Cnts]	<ul> <li>Active</li> </ul>
	Min 0 -	Max 1158
	Fast Lifetime[ns]	▼ Active
ster Carl		
	Min 0,000 -	Max 43,323
200 µm		

• Set the intensity scale from 0 counts – 200 counts and the lifetime scale from 0 to 5 ns.

FLIM Image Events[Cnts]	FLIM Image		
Events[Cnts]	FLIM Image		
		Events[Cnts]	
200 μm	<u>200 µm</u>	Min 0 Active Fast Lifetime[ns] Active Min 0,000 Max 4	

- In the image parameter panel, open the "Threshold" drop down menu.
- Activate "Use Threshold" and enter a threshold of 100.



## **Response:**

In the image, only pixels with higher photon counts are highlighted.

#### Note:

One can also use the bar in the intensity histogram to set the threshold.

FLIM Image		
FLIM Image		
	Events[Cnts]	
<u>200 μm</u>	✓ Min 0 ↔ Max 200 ↔ Fast Lifetime[ns] ✓ Active	
· · · · · · · · · · · · · · · · · · ·		

- For the analysis, only the spots within the nuclei of the cells are of importance. Therefore exclude the bright dots outside the nucleus of the cell on the lower right by placing the mouse cursor over the image, activate the context menu with a right mouse click, select "Rectangle ROI", press continuously "CRTL" and draw circles around the spots outside the nucleus.
- **Response:** The marked regions are excluded from analysis.

FLIM Image		
FLIM Image		
	Events[Cnts]	
	Min 0 Max 200 Fast Lifetime[ns]	
	Min 0,000 🖨 🔽 Max 5,000 🖨	
<u>200 μm</u>		

• In the lifetime fitting pannel, set the Fitting model to "Multi-Exp. Donor".

Fitting Model:	Multi-Exp. Donor
Decay:	Overall Decay
IRF: Import	Calculated IRF Remove
Model Parameters:	n 1 📥

• Click on IRF "Import"

Fitting	Model:	Multi-Exp. Donor	<b></b>	Help
Decay	:	Overall Decay	•	
IRF:	Import	Calculated IRF		Remove
Model	Parameters:	n	1 🜩	

• In the window appearing, select the file CENP-labelled cell\_for\_FRET\_IRF\_Det2.ptu as before in the FLIM script and click "OK".

🦗 Import IRF	
Samples	
Ey5_diff_IRF+FLCS-pattern.ptu سلاس	
Atto655_diff_FLCS-pattern.ptu سُلُسَتَ	
Atto655+Cy5_diff_FCS+FLCS.ptu لىلىس	
Atto655_diff_2FFCS.ptu لىلىس	
Atto488_diff_cw_total_correlation.ptu للملد	
Atto488_diff_pulsed_antibunching.ptu 🗄 🖓 🗄	
È Cy5_immo_FLIM+PoI-Imaging.ptu	
Lifetime_Trace.ptu سلس⊶ £ Cy5_immo_Lifetime_Trace.ptu	
Atto655_immo_On-Off-Analysis.ptu ليليس	
DaisyPollen_cells_FLIM.ptu	
GFP_RFP_cells_FLIM-FRET.ptu	
Gy3+Cy5_diff_PIE-FRET.ptu ليله Cy3+Cy5_diff_PIE-FRET.ptu	
TS-Bead_immo_xy-scan_Dual Focus.ptu	
TS-Bead_immo_xz-scan.ptu	
BA488+IBA547_unlinked_mix.ptu بالله BA488+IBA547_unlinked_mix.ptu	
BA488+547_crosslinked.ptu السلاس	
Atto488_485nm_pulsed.ptu المليس	
NV-Center_for_Antibunching_1.ptu سلاس	
NV-Center_for_Antibunching_2.ptu بالله NV-Center_for_Antibunching_2.ptu	
NV-Center_for_Antibunching_several.ptu	
CENP-labelled_cells_for_FRET.ptu	
CENP-labelled_cells_for_FRET_IRF_Det1.ptu للسلام	
CENP-labelled_cells_for_FRET_IRF_Det2.ptu	
FRET_GFP and mRFP.ptu	
Cancel	ок

• Activate the imported IRF by choosing it in the IRF option.

Fitting Model:		Multi-Exp. Donor	Help
Decay		Overall Decay	
IRF:	Import	Calculated IRF	Remove
Model	Parameters:	Calculated IRF CENP-labelled_cells_for_FRET_IRF_Det2.p	

**Response:**The imported IRF is highlighted in red.

Fitting Model:	Multi-Exp. Donor Help	Fit
Decay:	Overall Decay	Ν
IRF: Import	CENP-labelled_cells_for_FRET_IRF_De Remove	103
Model Parameters:	n 1 🔺	
Parameter	Value Fit 🔺	
Rol	1,000 🜩 📃	養 10 <sup>2</sup>
τ <b>ρ[ns]</b>	0,000 🗢	1
A[ŋ [kCnts] Limit	s 🗘 🗸	
τ[1] [ns] Limit	8 0,000 ± 0,000 € ✓	≝ 101
Bkgr Dec [Cnts] Limit	s 0,000 ± 0,000 € ✓	
Shift RF [ns] Limit	s 0,000 ± 0,000 € ✓	
Bkgr rer [Cnts] Limit	s 0,000 ± 0,000 € ✓	
EQ	0,000 ± 0,000	
E 🤹 [%]	0,000 ± 0,000	
R[	0,000 ± 0,000	Z 0,4
I[1] [kCnts]	0,000 ± 0,000	출 0,2
Isum[kCnts]	0,000 ± 0,000	<u>8</u> 0,0
A sum [kCnts]	0,000 ± 0,000	뿔 -0.2
τ Av int [NS]	0,000 ± 0,000	0 0 4 -
t Av Amp [NS]	0,000 ± 0,000	0 10 20 30 40 50
Clear	itiai Fit Fit X2 = 0,000	Time [ns]

• As model parameters, select n="2".

## Note:

If a mixture of complexes performing FRET and not performing FRET is present, the decay can of course be more complex, as the Cerulean alone already shows a bi-exponential decay behavior. However, the limited number of photons per pixel does not allow to use a more complex model.

- In the parameter panel, enter "2.675" ns as  $\tau_{\scriptscriptstyle D}$ 

## Note:

Whether you need to type "2.675" or "2,675" depends on the country specific settings of Windows program. For this tutorial, a computer with German specific settings was used, therefore, the decimal digits are separated with "," instead of ".".

Fitting Model:			/lulti-Exp. [	Donor				Не	lp	
Decay:			Overall Dec	cay			•			
IRF:	Impor	t – ] (	ENP-label	led_cells	_for_F	RET_IRF_De		Rem	ove	
Model Parameters:			n			2				
Parameter			Value						Fit	
Ro[]							1,0	00 🌲		
τo[ns]							2,6	75 🌲		
A [1] [kCn	ts]	Limits				0,000 ± 0,0	000	•	<	
A [2] [kCnts] Limits						0,000 ± 0,0	000	•	٨	
τ[1] [ns] Limits						0,000 ± 0,0	000	•	٨	
τ <sub>[2]</sub> [ns] Limits						0,000 ± 0,0	000	•	٨	
Bkgr Dec [	kgr Dec [Cnts] Limits					0,000 ± 0,0	000	•	<	
Shift IRF [	ns]	Limits	0,000 ± 0,000			•	>			
Bkgrirrf (	Cnts]	Limits				0,000 ± 0,0	000	•	>	
E[]						0,000 ± 0,0	000			
E % [%]						0,000 ± 0,0	000			
R[]			0,000 ± 0,000							
I[1] [kCnts]			0,000 ± 0,000							
I[2] [kCnts]						0,000 ± 0,0	000			
Isum[kCnts]						0,000 ± 0,0	000			-
Clea	ır	Ini	Initial Fit X <sup>2</sup> = 0,00			00				

• Click "Initial Fit".

**Response:**The fit is performed to the overall decay.

Fitting Model:	I	Multi-Exp. Donor Help	Fit
Decay:	(	Overall Decay	Λ
IRF: Import		CENP-labelled_cells_for_FRET_IRF_De	103
Model Parameters		n 2 🔺	
Parameter		Value Fit	
Roll		1,000 🜩 🗕	- Ê <sup>104</sup>
τ <b>ρ[ns]</b>		2,675 🚔	
A [1] [kCnts]	imits.	2,172 ± 0,000 🔶 🗸	
A [2] [kCnts]	imits.	1,906 ± 0,000 🔶 🗸	
τ[1] [NS] [	.imits	1,199 ± 0,000 🔶 🗸	
τ[2] [ns] [	.imits	3,481 ± 0,000 🔶 🗸	
Bkgr Dec [Cnts]	.imits	2,738 ± 0,000 🝨 🗸	
Shift ref [ns]	.imits	-0,216 ± 0,000 🝨 🗸	102
Bkgr IRF [Cnts]	.imits	0,264 ± 0,000 🔶 🗸	
ED		0,153 ± 0,000	
E 🤹 [%]		15,322 ± 0,000	
R[		1,330 ± 0,000	
I[1] [kCnts]		162,684 ± 0,000	مانى 🕹 🕹 🕹 🕹 مەلكە بولەر ئالىكە بارلىيە بەر 🕹 🕹 🗧 🗧
Ipg [kCnts]		414,562 ± 0,000	
Isum [kCnts]		577,246 ± 0,000	▼ -4 0 10 20 30 40 50
Clear	Ini	tial Fit X <sup>2</sup> = 1,040	Time [ns]

• In the Fitting panel, set the parameters  $Bkgr_{Dec}$ ,  $Shift_{IRF}$  and  $Bkgr_{IRF}$  constant by removing the mark. Set the background of the decay to 0.

## Note:

In order to reduce statistical fluctuations, as many parameters as possible need to be set constant. In a single pixel, the background counts of the pixel are negligible, therefore it is valid to set this to 0 in this case. Don't press "Initial Fit again".

Parameter	Value		
R <sub>0</sub> []	1,000		
τo[ns]	2,675	-	
A [1] [kCnts] Limit	2,172 ± 0,000	~	
A [2] [kCnts] Limit	1,906 ± 0,000	~	
τ[1] [ns] Limit	1,199 ± 0,000	~	
τ <sub>[2]</sub> [ns] Limit	3,481 ± 0,000	~	
Bkgr Dec [Cnts] Limit	p,000 ± 0,000		
Shift IRF [ns] Limit	-0,216 ± 0,000		
Bkgr IRF [Cnts] Limit	0,264 ± 0,000		

• Press "Calculate FRET" in the upper panel on the left.

#### Note:

Depending on the screen resolution, one may have to use the scroll-bar in order to access this button.



#### **Response:**

- A double exponential fit with the chosen start parameters is calculated for each selected pixel.
- From the results, a "FRET Image is calculated which is plotted in the FLIM image area. The FLIM image is hidden behind in another tab.
- On the right, the FRET efficiency histogram is plotted from the pixel values.
  - This graph shows 2 peaks, which is expected, as in the image one can see two cells with donor and acceptor construct and two cells without. For the cells with the FRET donor only, a FRET efficiency around 0 is expected and also shows up in the histogram.
- On the lower right, the FRET distance histogram is plotted. As we didn't enter any value of the Förster distance R0, the distance is plotted in terms of "R0".

## Note:

To transfer this in a nanometer scale, the real Förster distance must be known. The result is only meaningful, if the donor to acceptor ratio is 1:1 on the molecular level for each complex. This is not the case in our biological example with a more complex stoichiometry, therefore the graph is meaningless in this specific case.

Calculating the real Förster distance requires some effort and distinct knowledge about the sample. A few aids are available on the lab. These are not connected to PicoQuant and PicoQuant is not responsible neither for the content nor any possible errors; still these links may give an idea of what is required to calculate Förster distances.

- http://www.pymolwiki.org/index.php/Forster\_distance\_calculator
- http://www.osmanbilsel.net/software/forster-distance-calculator
- http://en.wikipedia.org/wiki/F%C3%B6rster\_resonance\_energy\_transfer



• In order to improve the FRET contrast, adjust the intensity scale to 0 counts - 200 counts.

#### Note:

The FRET efficiencies calculated also show values below 0. Although physically impossible, mathematically this reflects the variation in the fit.



• Press "Save Result" to save the current analysis.

File			
		Save Resi	ult
	Save Defaults		Restore Defaults

### **Response:**

A result file is saved and linked to the raw data file.



- The image analysis is now finished. There are several possibilities to continue:
  - To adapt the histogram plots, change the plot parameters on the right of the histogram to the desired values and click "Recalculate". To save these plots, click "Save Results" again.
  - To export certain analyses, place the mouse cursor over the area of interest, activate the context menu with a right mouse click and choose the export function of interest to export the data.
  - To analyze many images with the same FRET pair, it is possible to save the starting values for the fits by clicking on "Save Defaults" and store the starting values as a User Profile by entering the main menu ("Settings\Save User Profile as"). The software will need to restart to adapt the changes.
- If you don't want to do anything more, just close the result window.

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