# MANUAL

# Internally Quenched Fluorescence – Diubiquitin (IQF-DiUb) Substrates

### **Catalog Numbers:**

K11-linked Individual: DU1104

K48-linked Individual: DU4801, DU4802, DU4803, DU4804, DU4805, DU4806 Combination Panel: DU0101

K63-linked Individual: DU6301, DU6302, DU6303, DU6304, DU6305, DU6306 Combination Panel: DU0102

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# BACKGROUND

The ubiquitin-proteasome pathway plays a key role in protein modification and degradation. Ubiquitylation is a dynamic and reversible process mediated by the action of deubiquitylating enzymes (DUBs), a family of proteases that specifically cleave ubiquitin-derived substrates. The distinct structures of polyubiquitins determined to date suggest that DUBs act with high specificity towards various Ub linkage types (e.g. K48, K63, K11, etc). DUBs, in general, show remarkable diversity and the members of this family exhibit characteristic developmental and spatial expression patterns, as well as complex biochemical properties. The ability of DUBs to interact with preferred targets, thus modifying cellular functions, further highlights the importance of this pathway in both health and disease. Deregulation of ubiquitylation has been associated with a wide range of pathologies including cancer, muscle atrophy, infectious diseases and neurodegeneration and represents a promising target for therapeutic intervention. One of the challenges in monitoring DUBs activity is to create physiologically relevant substrate(s) to measure true isopeptidase activity in a format amenable to high-throughput screening (HTS). Currently, commercially available reagents and assays for measuring the activity of Ub/Ubl isopeptidases are based on the cleavage of a linear peptide- or amide-bond that do not reflect the geometry of a true isopeptide bond. In addition, other fluorescence based homogeneous assays for monitoring polyubiquitin chain disassembly suffer from the requirement of multiple enzymatic steps and/or N-terminal modification of ubiquitin molecule, a step homogenous, biologically relevant, and HTS amenable assay for DUBs.

### Diubiquitin (DiUb): A Novel Substrate for Robust Fluorescence Readout of DUBs Activity

LifeSensors has developed a line of novel physiological substrates for DUBs – diubiquitin molecules (Diubiquitin, DiUb) that are linked by isopeptide bonds via either K48 or K63, the most abundant forms of poly-ubiquitin linkages. These substrates can be used to determine the substrate specificity of numerous DUBs, to monitor the kinetic parameters of DUB mediated isopeptide cleavage, as well as investigate selective de-conjugation of poly-ubiquitylated proteins.



#### Assay versatility

There is a remarkable difference between the determined structures of K48- and K63-linked ubiquitin molecules, potentially providing the basis for their distinct functions. Moreover, the distinct structure of polyubiquitin linkages further illustrates a tremendous diversity among DUBs with regard to their substrate specificity. LifeSensors' IQF-DiUbs offer a variety of K48 and K68-linked substrates with FRET pair fluorophores uniquely positioned on specific sites of the diubiquitin molecule, thus providing a highly efficient tool to measure selective activity of the DUB of interest.

### About the assay

These diubiquitin substrates represent a new class of continuous assay substrates for the cleavage of a true isopeptide bond. The C-terminus of wild type ubiquitin is conjugated via an isopeptide bond to lysine 48 or lysine 63 of a second ubiquitin molecule, with the resultant diubiquitin forming an internally quenched fluorescence FRET pair (IQF) due to a presence of a highly efficient fluorescence quencher on one ubiquitin molecule and a fluorescent reporter (TAMRA) on the second ubiquitin molecule. Cleavage of the diubiquitin molecule by selective DUBs leads to separation of the fluorophore from quencher and subsequent increase in fluorescence signal. Each diubiquitin substrate is prepared through site specific labeling at different positions (K63-1-2-3-4-5-6 or K48-1-2-3-4-5-6) in order to allow optimization of a given diubiquitin substrate for an individual DUB. Importantly, the introduction of the fluorophore or quencher residue does not affect the native structure of the ubiquitin molecule. The generation of ubiquitin substrates represents a major advancement in the study of this important domain of the eukaryotic proteome.

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# **BENEFITS**

- IQF-DiUbs provide a sensitive, rapid, and robust fluorescent readout of enzymatic activity with minimal interference from screening compounds
- The assay measures cleavage of a physiologically relevant isopeptide bond rather than the linear peptide bond found in other currently available commercial assays for deubiquitylases
- A variety of DiUbs allows selection of the best substrate for your enzyme

## **APPLICATIONS**

- Investigation of the linkage specificity of DUBs
- Identification of agonists or antagonists of specific isopeptidases for HTS and drug discovery
- Determine kinetic parameters mediating substrate/DUB interactions

# **COMPONENTS**

### Reagents are supplied as individual DiUb substrates or as panels of smaller sizes

Buffer: 50mM Sodium MES, pH 6.0.

### Individual IQF Diubiquitins:

<b>K48-linked</b> 50μg @ 20μM each		<b>K63-linked</b> 50µg @ 20µM each						
Item Name (Cat.#)	Mol. Weight	Item Name (Cat.#)	Mol. Weight					
DiUb K48-1 (DU4801)	18,507.0	DiUb K63-1 (DU6301)	18,507.0					
DiUb K48-2 (DU4802)	18,548.4	DiUb K63-2 (DU6302)	18,548.4					
DiUb K48-3 (DU4803)	18,479.3	DiUb K63-3 (DU6303)	18,479.3					
DiUb K48-4 (DU4804)	18,520.4	DiUb K63-4 (DU6304)	18,520.4					
DiUb K48-5 (DU4805)	18,478.3	DiUb K63-5 (DU6305)	18,479.3					
DiUb K48-6 (DU4806)	18,520.4	DiUb K63-6 (DU6306)	18,520.4					

### Combination Panels of IQF Diubiquitins:

Item Name (Cat.#)	Description
DiUb K48 1-6 Panel (DU0101)	A sampler pack of six K48-linked IQF Diubiquitins: 25µL @ 20µM (~9.2µg) of each
DiUb K63 1-6 Panel (DU0102)	A sampler pack of six K63-linked IQF Diubiquitins: 25µL @ 20µM (~9.2µg) of each
DiUb K48 1-6 & DiUb K63 1-6 Panel (DU0201)	A sampler pack of six K48-linked and six K63-linked IQF Diubiquitins: 25µL @ 20µM (~9.2µg) of each

## ADDITIONAL ITEMS REQUIRED

- 1. Control isopeptidase: USP2 core (LifeSensors Cat. no DB501). USP2 core represents the common catalytic core domain of the two isoforms of USP2, USP2a and USP2b. This enzyme is known to cleave polyubiquitin chains formed through both K48 and K63 linkage types. As such, it is active against most of the IQF-DiUb substrates.
- 2. DUB of choice: LifeSensors provides the most comprehensive selection of DUBs available. Please visit <u>www.lifesensors.com</u> or contact a sales representative to learn more.
- **3.** Assay Buffer: 50mM Tris, pH 8.0, 0.05% CHAPS, 10mM DTT or buffer of choice. The addition of DTT or other reducing agent is required for the assay. Assay condition should be optimized by the end user.
- 4. 384-well black assay plates (Greiner BioONE 781209) or 96- well black assay plates (Greiner BioONE 655076).
- 5. Fluorescence plate reader. Filters or monochromators compatible with monitoring the fluorescence of TAMRA (Exc. 540 nm/Emm. 580 nm) are required. In addition, the use of a dichroic mirror with a cutoff in the range of 550-570 is highly recommended to ensure optimal signal-to-background. Further optimization of the plate reader optics (e.g. signal gain, plate height reads, etc.) is also recommended. Any fluorescence or multimode plate reader capable of the configuration described above should be suitable for this assay.
- 6. Ub-TAMRA Reference Standard (Cat.#DU0121). The standard can be used to optimize and calibrate the performance of individual plate readers (or fluorometers) for measurement of the rates of hydrolysis of the IQF-Diubiquitins by DUBs.

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# SUGGESTED PROTOCOL FOR MONITORING IQF-DIUb CLEAVAGE BY USP2 CORE

To obtain reliable and reproducible results, all samples should be run at least in duplicate. Allow all reagents to warm to room temperature (20-25°C) before use in the assay. **IQF-DIUD SUBSTRATES ARE LIGHT SENSITIVE AND MUST BE PROTECTED FROM LIGHT AT ALL THE TIMES** 

- 1. Dilute individual or combo panel DiUb substrates to 400nM, or 2x the desired final concentration, in assay buffer.
- 2. Prepare control USP2 core. Dilute USP2 core to 200 nM (2X) in assay buffer. A range of enzyme from 10nM to 1µM (final concentration) can be used for USP2 core control to generate a standard curve.
- 3. For 96-well plate, add 50 µL of USP2 core from step 2 to each control well of black assay plate (row G).
- 4. Prepare three dilutions of each DUB to be tested. (Suggested: 200nM, 100nM, 10nM).
- 5. Add 50 µL of assay buffer to "no enzyme" wells (row H).
- Dispense 50μL of each DiUb substrate from step 1 into each well. Note: the concentrations of both DiUb and DUB now is1/2 of the original (Steps 1 and 2).
- 7. Perform a kinetic read for 30min to 1hour. Important: The plate should be read immediately after the addition of the enzyme.

# WELL MAP

Typical 96-well assay set-up for either the DiUb panel (A) or a single IQF-DiUb (B) The number of DUBs of interest should be determined by the end user. Concentration of USP2 core used: 100 nM. Samples (DUB1 & DUB2) are loaded in duplicates (wells 1 and 2) in three dilutions (200nM, 100nM, 10nM).

Α.	<u>DU4801</u> <u>DU</u>		<u>4802 DU4</u> L I		<u>1803</u> <u>DU4</u>		<u>804</u> <u>DU4</u>		805 <u>DU48</u>		<u>806</u>	
	1	2	3	4	5	6	7	8	9	10	11	12
A	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1
	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM
В	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1
	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM
С	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1	DUB1
	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM
D	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2
	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM	200nM
E	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2
	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM	100nM
F	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2	DUB2
	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM	10nM
G	USP2	USP2	USP2	USP2	USP2	USP2	USP2	USP2	USP2	USP2	USP2	USP2
	100	100	100	100	100	100	100	100	100	100	100	100
	nM	nM	nM	nM	nM	nM	nM	nM	nM	nM	nM	nM
Н	No	No	No	No	No	No	No	No	No	No	No	No
	enzyme	enzyme	enzyme	enzyme	enzyme	enzyme	enzyme	enzyme	enzyme	enzyme	enzyme	enzyme

**B.** DUB screening against a single substrate. DUBs of interest are loaded in triplicates. DiUb substrates should be prepared in two concentrations (400nM and 200nM, <u>2X</u>). Final concentration is shown on the left.

		DUB1			DUB2			DUB3			DUB4		
DiUb	A	DUB1	DUB1	DUB1	DUB2	DUB2	DUB2	DUB3	DUB3	DUB3	DUB4	DUB4	DUB4
subst.		200nM											
100	В	DUB1	DUB1	DUB1	DUB2	DUB2	DUB2	DUB3	DUB3	DUB3	DUB4	DUB4	DUB4
nM		100nM											
100	С	DUB1	DUB1	DUB1	DUB2	DUB2	DUB2	DUB3	DUB3	DUB3	DUB4	DUB4	DUB4
nM		10nM											
100	D	DUB1	DUB1	DUB1	DUB2	DUB2	DUB2	DUB3	DUB3	DUB3	DUB4	DUB4	DUB4
nM		200nM											
200	Е	DUB1	DUB1	DUB1	DUB2	DUB2	DUB2	DUB3	DUB3	DUB3	DUB4	DUB4	DUB4
nM		100nM											
200	F	DUB1	DUB1	DUB1	DUB2	DUB2	DUB2	DUB3	DUB3	DUB3	DUB4	DUB4	DUB4
nM		10nM											
200 nM	G	USP2 100 nM											
	Н	No enzyme											

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# **STORAGE**

IQF-DiUbs can be stored at 4°C for up to 6 months. Long term (> 6 months) storage at -80°C is recommended. If the products are received frozen on dry-ice please store at -80°C until ready to use. Avoid repeated freeze/thaw cycles. **IQF-DiUb substrates are light sensitive and must be protected from light at all times.** 

Please note that some physical characteristics and protocols are item specific. Please refer to individual product sheets or application notes now available at <u>www.lifesensors.com</u> for further information.

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