

Sep 28, 2018

IRDye® Peptide Labeling Application Guide 🖘

LI-COR Biosciences1

¹LI-COR Biosciences

LI-COR Biosciences

dx.doi.org/10.17504/protocols.io.guwbwxe

Working



Margaret Dentlinger 🕜 🦰 😱





ABSTRACT

IRDye NHS esters react with unblocked amines in peptides. NHS ester reactions usually proceed quickly and cleanly in either organic or aqueous solvents. While NHS esters continue to be widely used tools for biomolecule modification, their application to large peptides may be complicated by factors such as steric hindrance, amine inactivation by salt-bridging, and multiple labeling sites.

Although a properly designed peptide can be labeled site-specifically by NHS ester chemistry, complex designs may incur costprohibitive materials and labor. To address the limitations of NHS ester chemistry, orthogonal technologies such as IRDye Maleimides and Click Chemistry Reagents have been developed to label non-amine functional groups. Orthogonal chemistry can also facilitate dual- or multi-dye labeling. As with NHS ester chemistry, optimal non-amine labeling is usually achieved by aqueous solution-phase reactions.

Developed for:

Odyssey® Classic, Odyssey CLx, Odyssey Sa, and Aerius Imaging Systems

EXTERNAL LINK

https://www.licor.com/documents/nmekjs7iez6sw5p8fv7b7005chbrcog7

AppNote PeptideLabe ling_0117_988-13575.pdf

PROTOCOL STATUS

Working

We use this protocol in our group and it is working

GUIDELINES

I. Introduction

Peptides labeled with fluorescent dyes are important as probes for in vivo imaging and as substrates for enzyme activity assays. Near-infrared (NIR) fluorophores such as IRDye 800CW (excitation 780 nm; emission 794 nm) can offer improved sensitivity because of low NIR autofluorescence from tissues, cells, biological materials, or drug compounds. Fluorophores can also be combined with appropriate quencher dyes to create fluorogenic peptide probes, such as those used for protease activity detection. 1-4 IRDye QC-1 can efficiently quench a wide range of fluorophores spanning the visible and NIR spectra (~500 - 800 nm) in a fluorescence reso- nance energy transfer (FRET) system. Together, IRDye 800CW and IRDye QC-1 comprise an optimal fluorophore-quencher pair for incorporation into NIR fluorogenic peptides (Figure 1).

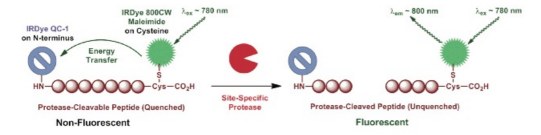


Figure 1. NIR fluorogenic peptide probe using IRDye 800CW donor and IRDye QC-1 quencher.

IRDye fluorophores and quenchers are amenable to a wide variety of conjugation reactions that furnish stable, covalently-labeled peptides. Furthermore, site-specific dye labeling can be achieved by employing orthogonally reactive groups in peptide design. Because IRDye labels have been optimized for detection in aqueous biological environments, conjugation reactions are best performed in solution-phase on fully deprotected peptides.

II. Peptide Labeling Options

IRDye NHS esters react with unblocked amines in peptides. NHS ester reactions usually proceed quickly and cleanly in either organic or aqueous solvents. While NHS esters continue to be widely used tools for biomolecule modification, their application to large peptides may be complicated by factors such as steric hindrance, amine inactivation by salt-bridging, and multiple labeling sites.

Although a properly designed peptide can be labeled site-specifically by NHS ester chemistry, complex designs may incur cost-prohibitive materials and labor. To address the limitations of NHS ester chemistry, orthogonal technologies such as IRDye Maleimides and Click Chemistry Reagents have been developed to label non-amine functional groups. Orthogonal chemistry can also facilitate dual- or multi-dye labeling. As with NHS ester chemistry, optimal non-amine labeling is usually achieved by aqueous solution-phase reactions. Table 1 summarizes general strategies for labeling peptides with IRDye reagents.

Table 1. IRDye Reagents for Labeling Peptides

Desired Peptide	Appropriate	Notes
Labeling Site	IRDye Reagent	
Unblocked N-terminus or	NHS Ester	Perform in mildly basic aqueous solutions; peptides
Side-Chain Amine (e.g., lysine)		containing >1 amine may yield product mixtures
Sulfhydryl (e.g., cysteine)	Maleimide	Perform in mildly acidic aqueous solutions
		for selective labeling in the presence of amines
Alkyne	Azide	Copper-catalyzed Click Chemistry for selective
		labeling in the presence of amines
	Alkyne	Copper-catalyzed Click Chemistry for selective
Azide		labeling in the presence of amines
DBCO	Copper-free Click Chemistry for selective	
	labeling in the presence of amines	
DBCO	Azide	Copper-free Click Chemistry for selective labeling
		in the presence of amines

IRDye Maleimides are designed to react selectively with sulfhydryl groups in the presence of amines.

Optimal selectivity is usually obtained with labeling reactions performed in mildly acidic (pH 6.5) aqueous solvents; at acidic pH, amines are rendered inert by protonation. Sulfhydryl groups can be installed in peptides with cysteine or commercially-available thiollinkers.

NOTE: Peptides containing sulfhydryls can spontaneously form inactive dimers. Therefore, it may be necessary to pre-treat the peptide with a reducing agent such as tris(2-carboxyethyl) phosphine (TCEP) to reveal active monomers.

IRDye Click Chemistry reagents are mostly inert to naturally-occurring functional groups such as amines and hydroxyls. Click Chemistry utilizes high-fidelity pairs of functional groups that can react either in the presence⁵ or absence⁶ of a copper catalyst. Click Chemistry functional groups can also be readily installed in peptides with commercially-available reagents.

III. Labeling Reaction Techniques

a. Aqueous Solution-Phase Labeling

For all labeling reactions, it is critical to omit components that can interfere with the coupling of the peptide and the dye. Peptides should be dissolved in aqueous buffers that do not contain extraneous nucleophiles such as Tris, sodium azide, DTT, BME, etc.To maximize the labeling efficiency, the peptide should be dissolved at the highest practical concentration that still maintains a homogeneous reaction mixture. The aqueous reaction mixture can be directly purified by reverse-phase HPLC (see Section III.C. Purification). Table 2 provides general parameters for aqueous solution-phase labeling of peptides with IRDye reagents.

Table 2. General parameters for labeling peptides in aqueous solutions

IRDye	Dye Equivalents	Reaction	Reaction	Reaction
Reagent	per Labeling Site	Buffer (pH)	Temperature	Time (hours)
NHS Ester	1-2	Phosphate	Ambient	2-3
		(8.0-8.5)	to 37 °C	
Maleimide	1-2	Phosphate	Ambient	2-3
		(6.5-7.5)	to 37 °C	
Alkyne†	1	Variable	Ambient	1-2
		(6.5 to 8.5)	to 37 °C	
Azide†,*	1	Variable	Ambient	1-2
		(6.5 to 8.5)	to 37 °C	
DBCO*	1	Variable	Ambient	1-2
		(6.5 to 8.5)	to 37 °C	

^{*} Copper-catalyzed Click Chemistry⁵

b. Organic Solution-Phase Labeling

Certain peptides (e.g., comprised mainly of hydrophobic amino acids) may not be amenable to aqueous solution-phase labeling. In these atypical cases, IRDye reagents can be used for organic solution-phase labeling; however, this reaction may be slower than its aqueous counterpart. Again, peptides should be dissolved in solvents devoid of extraneous nucleophiles such as Tris, sodium azide, DTT, BME, etc.The optimal reaction solvent is anhydrous dimethyl sulfoxide (DMSO). When labeling with IRDye NHS esters and Maleimides, including a tertiary amine base such as N,N-diisopropylethylamine (DIPEA) is necessary to promote the reaction. After the reaction has completed, the crude IRDye labeled peptide can be precipitated from the DMSO by dropwise addition of anhydrous diethyl ether. IRDye labels impart hydrophilicity and can facilitate the purification of hydrophobic peptides by reversephase HPLC (see Section III.c. Purification). Table 3 provides general parameters for organic solution-phase labeling of peptides with IRDye reagents.

Table 3. General parameters for labeling peptides in organic solutions

IRDye Reagent	Dye Equivalents per Labeling Site	Base Equivalents per Labeling Site	Reaction Temperature	Reaction Time (hours)
NHS Ester	1	2 - 4	Ambient to 37 °C	2 - 12
Maleimide	1	2 - 4	Ambient to 37 °C	2 - 12

[†] Copper-free Click Chemistry⁶

Alkyne†	1		Ambient	1 - 4
		None	to 37 °C	
Azide†,*	1		Ambient	1 - 4
		None	to 37 °C	
DBCO*	1		Ambient	1 - 4
		None	to 37 °C	

^{*} Copper-catalyzed Click Chemistry⁵

For questions, please send a detailed inquiry to biohelp@licor.com

c. Purification

Reverse-phase HPLC purification delivers IRDye dye-labeled peptides with the highest purity. Zorbax C18 HPLC columns (Agilent Technologies) work well, and are available in different sizes for analytical and preparatory scale. The recommended mobile phase is a gradient mixture of acetonitrile and water buffered with triethylammonium acetate (TEAA, 50 mM, pH 6.0). The TEAA provides ion-pairing for better retention behavior of the hydrophilic IRDye dye-labeled peptides on the column. As a consequence, however, the predominant counter-ion will be triethylammonium, which may interfere with downstream biological experiments. Prior to lyophilization, the purified peptide should be ion-exchanged by eluting through an appropriate resin (e.g., Amberlite $^{\text{m}}$) or by dialysis. For flexibility in development and/or troubleshooting, the HPLC system should be equipped with a diode array detector (DAAD) and be able to monitor at the absorption maxima of all dyes used.

For questions, please send a detailed inquiry to biohelp@licor.com

d. Quantification

UV-Vis absorbance spectroscopy is the best method for quantifying the amount of purified IRDye labeled peptide. To determine the concentration of a stock solution containing a singly-labeled peptide, dilute an aliquot of the stock solution in 1X PBS, measure the absorbance spectra at the dye-specific maximum (baseline subtraction from 950 to 1000 nm), and use the following equation:

Conc of IRDye peptide (mg/mL) =
$$\frac{A_{dye}}{\epsilon_{dye}}$$
 x Dilution Factor x MW_{IRDye peptide}

In which:

 $A_{\mbox{\scriptsize dye}}$ is the measured absorbance at the dye-specific maximum (See Table 4)

 ϵ_{dye} is the extinction coefficient for the dye in 1X PBS (See Table 4) Dilution Factor is the fold dilution of the IRDye peptide in 1X PBS

 $\ensuremath{\mathsf{MW}_{\mathsf{IRDye}}}\xspace_{\mathsf{peptide}}$ is the molecular weight of the IRDye peptide

Table 4. Parameters for UV-Vis quantification of IRDye labeled peptides in 1X PBS

Label	Dye Maximum (nm) for Measuring Absorbance	e dye
IRDye 800CW	774	240,000
IRDye 800RS	767	200,000

[†] Copper-free Click Chemistry⁶

IRDye 750	756	260,000
IRDye 700DX	689	165.000
IRDye 680RD	672	165,000
IRDye 680LT	676	250,000
IRDye 650	648	230,000
IRDye QC-1	737	96,000

For a pure stock solution of a FRET-quenched peptide labeled with exactly one IRDye 800CW and one IRDye QC-1: Determine the concentration of a stock solution by diluting an aliquot of the stock solution in methanol, measuring the absorbance spectra at 778 nm and 850 nm (baseline subtraction from 950 to 1000 nm), and using the following equation:

Conc of IRDye 800CW QC1 peptide (mg/mL) = $\frac{A_{778} - (1.265 \times A_{850}) \times D}{300.000}$ Dilution Factor × MW IRDye 800CW QC1 peptide

In which:

 $\ensuremath{A_{778}}$ is the measured absorbance at the maximum for IRDye 800CW in methanol

300,000 is the extinction coefficient for IRDye 800CW in methanol

1.265 is the correction factor for the ratio of A_{778/A850} for IRDye QC-1 in methanol

 A_{850} is the measured absorbance at the maximum for IRDye QC-1 in methanol

Dilution Factor is the fold dilution of the IRDye 800CW QC1-peptide in methanol

 $MWI_{RDye\ 800CW\ QC1\ peptide}$ is the molecular weight of the IRDye 800CW QC1 peptide

For questions, please send a detailed inquiry to biohelp@licor.com

IV. Examples

For an example of aqueous solution-phase labeling with IRDye 800CW NHS ester, see: Davies-Venn CA, Angermiller B, Wilganowski N, Ghosh P, Harvey BR, Wu G, Kwon S, Aldrich MB, Sevick-Muraca EM. Albumin-Binding Domain Conjugate for Near-Infrared Fluorescence Lymphatic Imaging. Mol. Imaging Biol. 14(2012), 301 – 314.

For an example of organic solution-phase labeling with IRDye 800CW NHS ester, see: Chen Y, Dhara S, Banerjee SR, Byun Y, Pullambhatla M, Mease RC, Pomper M. A low molecular weight PSMA-based fluorescent imaging agent for cancer. Biochem. Biophys. Res. Commun. 390(2009), 624–629.

For an example of aqueous solution-phase labeling with IRDye 800CW Maleimide, see: YeY, Zhu L, MaY, Niu G, Chen X. Synthesis and evaluation of new iRGD peptide analogs for tumor optical imaging. Bioorg. Med. Chem. Lett. 21(2011), 1146–1150.

For examples of dual-labeling with IRDye® QC-1 NHS ester and various fluorophores, see: Sun X, Zhang A, Baker B, Sun L, Howard A, Buswell J, Maurel D, Masharina A, Johnsson K, Noren CJ, Xu M-Q, Corrêa IR. Development of SNAP-Tag Fluorogenic Probes for Wash-Free Fluorescence Imaging. ChemBioChem 12 (2011), 2217–2226.

V. Additional References

Matayoshi ED, Wang GT, Krafft GA, Erickson J. Novel fluorogenic substrates for assaying retroviral proteases by resonance energy transfer. Science 247(1990), 954-958.

Bullok KE, Maxwell D, Kesarwala AH, Gammon S, Prior JL, Snow M, Stanley S, Piwnica-Worms D. Biochemical and in vivo characterization of a small, membrane- permeant, caspase-activatable far-red fluorescent peptide for imaging apoptosis. Biochem. 46 (2007), 4055-4065.

Zheng G, Chen J, Stefflova K, Jarvi M, Li H, Wilson BC. Photodynamic molecular beacon as an activatable photosensitizer based on protease controlled singlet oxygen quenching and activation. Proc. Natl. Acad. Sci. USA 104 (2007), 8989-8994.

Blum G, Mullins SR, Keren K, Fonovic M, Jedeszko C, Rice MJ, Sloane BF, Bogyo M. Dynamic imaging of protease activity with fluorescently quenched activity-based probes. Nature Chem. Biol. 1(2005), 203-209.

 $Best\ MD.\ Click\ Chemistry\ and\ Bioorthogonal\ Reactions: Unprecedented\ Selectivity\ in\ the\ Labeling\ of\ Biological\ Molecules.$ $Biochemistry\ 48(2009),\ 6571-6584.$

Simon M, Zangemeister-Wittke U, Plückthun A. Facile Double-Functionalization of Designed Ankyrin Repeat Proteins using Click and Thiol Chemistries. Bioconjugate Chem. 23(2012), 279-286.

VI. Associated Products

http://www.licor.com/bio/products/reagents/nhs_esters.jsp

http://www.licor.com/bio/products/reagents/maleimides/maleimides.jsp

http://www.licor.com/bio/products/reagents/click_chemistry/clickChemistryOverview.jsp

http://www.licor.com/bio/products/reagents/fret_based_assays/fret_based_assays.jsp

http://www.licor.com/bio/services/custom_labeling.jsp

http://www.licor.com/bio/network/#crb

Adheous	Solution-	Phase	Laheling

1 Dissolve peptides in aqueous buffers that do not contain extraneous nucleophiles such asTris, sodium azide, DTT, BME, etc.

NOTE

To maximize the labeling efficiency, the peptide should be dissolved at the highest practical concentration that still maintains a homogeneous reaction mixture.

For all labeling reactions, it is critical to omit components that can interfere with the coupling of the peptide and the dye.

9 Purify the aqueous reaction mixture directly by reverse-phase HPLC (Purification-Step 6).

See Table 2. in Guidelines for general parameters for aqueous solution-phase labeling of peptides with IRDye reagents.

Organic Solution-Phase Labeling

- Certain peptides (e.g., comprised mainly of hydrophobic amino acids) may not be amenable to aqueous solution-phase labeling. In these atypical cases, IRDye reagents can be used for organic solution-phase labeling; however, this reaction may be slower than its aqueous counterpart.
 - Again, dissolve peptides in solvents devoid of extraneous nucleophiles such as Tris, sodium azide, DTT, BME, etc.
 - The optimal reaction solvent is anhydrous dimethyl sulfoxide (DMSO).

NOTE

When labeling with IRDye NHS esters and Maleimides, including a tertiary amine base such as N,N-diisopropylethylamine (DIPEA) is necessary to promote the reaction.

- 4 After the reaction has completed, precipitate the crude IRDye labeled peptide from the DMSO by dropwise addition of anhydrous diethyl ether.
- IRDye labels impart hydrophilicity and can facilitate the purification of hydrophobic peptides by reverse phase HPLC (Purification-Step 6).
 - See Table 3 in <u>Guidelines</u> for general parameters for organic solution-phase labeling of peptides with IRDye reagents.

NOTE

For questions, please send a detailed inquiry to biohelp@licor.com

Purification

- 6 Reverse-phase HPLC purification delivers IRDye dye-labeled peptides with the highest purity. Zorbax C18 HPLC columns (AgilentTechnologies) work well, and are available in different sizes for analytical and preparatory scale.
 - The recommended mobile phase is a gradient mixture of acetonitrile and water buffered with triethylammonium acetate (TEAA, 50 mM, pH 6.0).*As a consequence, the predominant counter-ion will be triethylammonium, which may interfere with downstream biological experiments.
 - Prior to lyophilization, the purified peptide should be ion-exchanged by eluting through an appropriate resin (e.g., Amberlite ™) or by dialysis.

NOTE

For flexibility in development and/or troubleshooting, the HPLC system should be equipped with a diode array detector (DAAD) and be able to monitor at the absorption maxima of all dyes used.

Quantification

7 UV-Vis absorbance spectroscopy is the best method for quantifying the amount of purified IRDye labeled peptide. To determine the concentration of a stock solution containing a singly-labeled peptide, dilute an aliquot of the stock solution in 1X PBS, measure the absorbance spectra at the dye-specific maximum (baseline subtraction from 950 to 1000 nm), and use the following equation:

Conc of IRDye peptide (mg/mL) = $\frac{A_{dye}}{\epsilon_{dye}}x$ Dilution Factor x MW_{IRDye peptide}

In which:

- Adye is the measured absorbance at the dye-specific maximum (See Table 4)
- ε dye is the extinction coefficient for the dye in 1X PBS (SeeTable 4)
- Dilution Factor is the fold dilution of the IRDye peptide in 1X PBS
- MWIRDye peptide is the molecular weight of the IRDye peptide

This is an open access protocol distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited