Abstract

Program # 502.17, Abstract # 6822 Reliable duplex and multiplex assays remain a challenge for fluorescence reporter enzyme technology. To this end we aveloped a system that takes advantage of pro-fluorescent es and a novel azoreductase.

mized and used to bacterially express the protein. Biochemical acterization of the purified enzyme in solution revealed a mer carrying a non-covalently bound flavin mononucleotide and the set of the se dride transfer reaction were consistent with the bi-bi ping-pon-achanism of related oxidoreductases. In our optimized colorimetri an that of β-galactosidase.

ontaining BHQs and a range of fluorophores readily released uorescence in an NADPH-dependent manner with robust signal to

In short, by matching BTI azoreductase with pro-fluorescent trates we have created a simple, yet adaptable, reporter system

The BTI1 Azoreductase Colorimetric and Fluorometric Reporter System

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Background and Goals

Reporter enzyme assays are employed to assess the effect of cellular stimuli with high sensitivity. The preferred signal output is highly amplified, long lasting and free from background noise. Enzymes that release fluorescent signals are well suited for reporter amplification. A persistent limitation lies in the need for a compatible internal control enzyme assay. To complement existing fluorogenic assays with limited wavelength readouts, we set out to construct a reporter system that yields fluorescent signals over the visible spectrum.

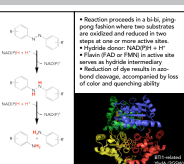
There are >10,000 man-made aromatic color fast azodyes used for textile, food and hair coloring. Azo dyes have broad absorption spectra with no fluorescence and are ideally suited as quencher dyes. With this in mind, Biosearch Technologies has developed a series of Black Hole Quenchers (BHQs) with a polyaromatic azo-backbone that have found widespread use in multiplex quantitative cDNA and genomic DNA PCR assays.

Azo dyes present an environmental burden in locals heavy with textile industries. To address this issue much research has gone into bacterial and fungal decolorization. Out of this research bacterial NAD(P)H-azoreductases have been isolated and characterized.

BHQs with different absorption ranges can readily be linked to fluorophores throughout the spectrum. The azoreductase reporter enzyme acting on such dark profluorescent substrates can therefore be combined with other fluorogenic reporter assays.

Here is described the procurement and biochemical characterization of the BTI1 azoreductase and our efforts of assembling a novel reporter enzyme system.

Enzymatic azoreduction



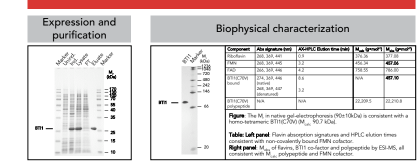
Materials and Methods

Cloning and expression The BTI1 azoreductase clone SO1-2 was isolated from a local soil sample by use of degenerate PCR. The azoreductase (C70V) open in the contract of the contrac

immunorecognition (myc) and affinity purification (His), Tagged BTI (C70V) was expressed and purified making use of the C-term His, tag from E. coli BL21(DE3) pL25 sessentially as recommended (Novagen). Routinely, 50 m cultures yielded 2.5 mg of 97% pure protein. Gel-electrophoresis Samples for SDS-PAGE were denatured in Learnmi loading buffer and separated on a 14% "IrScJy gel (Invitrogen). Samples in Tris-Cl gylcard) were separated on a 14% "Is native polyacyralmide gel (Invitrogen). Proteins in the gels were stained with GelCode (Prece). The size markers were: PAGE Ruler (Fermentas), Native-Mark (Invitrogen). PHCL and Mass Spectrometry The absorption spectra and the elution profiles in anion exchange HPLC of native and denatured BTI11(C70V) as well as of standard flavin co-factors were recorded on an HP diode array spectrometer and a Vaters HPLC. BTI11(C70) was subjected to ESI-MS (Waters) to determine Profiloarcent substrate synthesis and purification Profileson problems proteins of solosarch 8700 DNA synthesizer using standard phosphoramidite reagents. Samples were purified by XX-HPLX, and salts removed on a reverse phase cartridge. Purity was assessed by RP- and AX-HPLC. The absorption form free amines (Quencher deroder) produces in produces to CFI.

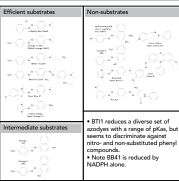
The absence of absorption from free amines (quencher degradation products) confirmed dye integrity. Quantitative azoreduction All buffers were at 200 mM and NADPH at 200 µM. In both absorbance and fluorescence assays, 200 µl samples were placed in 96-well plates. Typically 0.5-2.0 µg of enzyme was used per reaction. Apparent K_M and V_{max} values were determined by varying the concentration of azo dye substrates (0.5-50 µM) and making Lineweaver-Burk plots. Reaction rates were calculated by fitting the initial part of the absorbance decay by linear regression to find the slope. Absorption spectra were recorded in a HP 8453 diode array spectrophotometer, and absorbance measurements were recorded in a Tecan Safire plate reader

Results

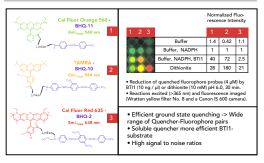


Enzymology Test substrates: Methyl Orange (MeO; Orange III); Water soluble pH indicator; pK_a 3.7; A_{max} (pH > 3.7) 466 nm (ɛ: 35900 M⁻¹•cm⁻¹) Flame Orange (Basic Orange 31, BO31); Water soluble hair dye (Ciba); pK_a > 8.5; A_{max} (pH < 8.3) 483 nm (ε: 28360 M⁻¹•cm⁻¹) Α A pH-profile of BTI1-mediated reduction of MeO (Black Open circles) (optimum 6.0), and FO (Red Open circles) (optimum > 8.3) B (Open circles) Effective temperature range for consistent azoreduction (at pH 6.0): 25-40 °C. (Open squares) Effective thermostability range <20-45 °C pH optimum substrate- and enzyme- dependent 0-mge NADPH preferred hydride donor (not shown) Onange II (OR) Mono- or di-valent cations not required (not shown) Tolerates moderate concentrations of cell extraction detergents and rganic solvents (not shown)

Substrate specificity



Fluorescence release



Summary

• The BTI1 azoreductase is a homo-tetrameric flavo-enzyme that uses a non-covalently attached FMN

Black Hole Quenchers and other azodyes are efficiently reduced by BTI1

over wide pH and temperature ranges • BTI1 releases fluorescent signals with high signal to noise ratios

• The enzyme's broad temperature optimum and and ability to reduce substrates with different pKas and absorption maxima will enable duplex and multiplex assays with reporter enzymes like β-galactosidase and alkaline phosphatase

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Notes

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