

# **Comparison of Activity of Botulinum Neurotoxin Type A Holotoxin and Light Chain Using SNAPtide<sup>®</sup> FRET Substrates**

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## Abstract

The brailmum neurotronism are composed of two submits, at 100 kD heavy chain and 5 ±0 kD light chain, which are litted by single dualified board. The heavy chain is representishe for braining and translocation while the light chain ontonics one tansumaria carivity. The light chain is a zine dependent metallegrosens. The enaltyre of washing metastratic for braining manetarious in type A is the synghtosamal protein SNAP-25. A previously reported, we have developed fluencences resonance energy transfer (RFR) business based on SNAP-25 that are reality resonation and light chain. One of the substrates contains an oAux DNP FRET pair and the other a HTCD DADCUC HERF pair (US spacest RNMOA).

In this such, the specific activity of the holotonic and recombinant light chain were determined using both PRIT substances. Specific activity is expressed in terms of purpose 35 SMN the<sup>24</sup> cleared pert minute perturbation of enzyme. For comparison, the specific activity of holotonic and light chain were accertained undergradiance distributions. The specific activity was also calculated for the light chain moder in optimal conditions for the specific concentrations were treaded under were also than used the optimal conditions. A strete of 2A/C, concentrations were treaded in several distributions of the strete distributions of the strete when the strete stre

## **Introduction**

The bothdium neurotoxis as one of the most point toxis is nature. Their postery, is human is due to the clearge and microsition of operform formula points. The temposable for the shading and the temposable structure of the structure of the structure of the shading and domain. The light-damain is the response perform of the temposable for the shading and domain. The light-damain is the response perform of the temposable for the shading and domain. The light-damain is the response perform the temposable temposable temposable which cleares a a specific site in one of the synghtsomal proteins. These as between a SNAB442, the light synthesis in for building neuroscient type A. These submissing are known as SNAB442, interlay excelled finate structure and the structure is the structure of the structure is the structure of the structur

Several of the light chains of the boulinum scenatorian have been expressed and particult as foriantion of the light chains of the boulinum scenatorian have been expressed and particult as foriant are unable to penetrate cells and cause toxisity. They represent an advertarity means of malying boulinum scenatorians without weaving algood the high bouxing and RL Conditions required and weaworking with the holotoxin. Here we compare the enzymatic activity of the full bouilinum neurostants (PA) (RTA) and the holotoxin. Here we compare the enzymatic activity of the full bouilinum neurostant (PA) (RTA) and the holotoxin. Here we compare the enzymatic activity of the full bouilinum neurostant activity and the holotoxin. Here we compare the enzymatic activity of the full bouilinum neurostant activity and the holotoxin. Here we compare the enzymatic activity of the full bouilinum neurostant activity and the holotoxin. Here we compare the enzymatic activity of the full bound of the activity and the holotoxin. Here we compare the enzymatic activity of the full bound enzymatic activity of the full bound of the strate are neurosed using the SNAP04<sup>4</sup> thorogenetic enzymatic activity and the observation.



SNAbide containing either the so-anisobenenic wide 24 distinguishes) (AshcDNP) FRIT pair or the merscenic thiochumbourgo/DARCT, (FIFT David to be teen synshesized TRA of LA enzymatic activity can be monitored continuously by recording the intrase in fluences in themsity over the FITCDARCT, abstrate and up to 30 dA for the oAShCPM substrate. An 8 fold increase is FITCDARCT, abstrate and up to 30 dA for the oAShCPM substrate. An 8 fold increase in fluencescone intensity over the oAShCPM substrate. An 8 fold increase in fluencescone intensity to sobered when the oAShCPM substrate. An 8 fold increase in the optimum strate and up to 30 dA for the oAShCPM substrate. The MCMCPL substrate and genera that 90 fold increase in fluencescone intensity to downeed for the TITCDARCYL substrate "Bottom".

# Materials

SNAPide<sup>®</sup> substrates (product #520 and #521), SNAPide<sup>®</sup> Unquenched Calibration Peptides (product #528 and #529), bolulium meurotoxin type A (product #130A), botulium meurotoxin type A light chain, recombinant (product #610A), bolulium metrovoxin type B light chain, recombinant (product #620A) and VAMPide<sup>®</sup> (product # 540) are all products of List Biological Laboratories, Inc.

## Methods

Flavariantic surger: Cominsons aways were performed on a SPECTRAms GEMNIX SS fluence-conce imorplate reader Molecular Devices, Smarphue CA, biai and Genet HU, DTRAC hale. In devicement plants: (EdS Scientific, Campbell, CA), Steck solutions of each FRET solutions were made in dimethyl auditodie DMSOS (Trail diditionis were made in the assays hifter, "Takes were include and 37°C for 15, minutes to ensure equilibrium prior to addition of substrate. The reaction was initiated by the addition of 37°C SAMIde<sup>2</sup> or VAMIde<sup>2</sup>." The time-dependent increase in fluencesce intensity was monitored at 37°C. The excitation and emission wavelengths were set to 321 nm and 418 nm, respectively, for the coher DMPstortmet and 490 nm d52 nm using a comfilter of 95 nm for the FTICDARCYL-Mortent. Zane threation: LeAs is the rise dependent endoperdates proting of balantam neuronizating etc. A During particulation of the high chain advantami, it is possible that the conductant zien modeluse may be loss from some of the postim molecules. To set if zinc could be added back is the enzyme and increase its enzymain activity, zince chiefuide was titeaded into several reaction HIPSES, pdf 18.0, 2014 HIPSES, pdf 18.0 - 0005 HIPSES, pdf 18.0 - 1000 HIPSES, pdf 18.0, 2014, was added to reactions staring as 4 and and intringing down to 0.001 and N = 21dd distinos. The reactions were monitored continuously for one how at 27 Agences Chiosin meaning BET (2014) enzyme in the 21dd distinos.

Later of Detection (LDD): The limit of directions is the minimum concentration of PLA or LeA that case how measured with WPA confidence that the concentration of RTA At a LA present is praret than zero under conconditions sing this flowappencie assay. A calibration curve was generated by measuring the initial rate of protochysis as a function of earyne concentration. Server policitas of a concentration of RTA At a LA present is praret than zero more than considered to the concentration. Server policitas of a concentration of RTA at LA present is praret than the concentration of RTA at LA and generation Distribution with WPA confidence and A degrees of freedom limits was calculated as 2.988 (Saleet's 6 Distribution with WPA confidence and A degrees freedom limits was any was also done with LA using coress were performed with RTA and LA using both SNAPaide's aburnture at 8 jukt. The assay was reactioned on at 13 degrees Calibration for alth M organs. The savey were carried on at 13 degrees Calibration for the and concentration of 3 juht (Hgrees 7). The savey were carried on at 13 degrees Calibration at the MPA at LA using coress.

VLMTMeV : VLMTMeV is a sew substate designed for discretising the activity of bondhum networking type II. To measure the ullity of diss substates is two networking informations mercoroticity per light chalm. (dd.). E. Left substates is a structure light polarization mercoroticity per light chalm. (dd.). E. Left per light charge of the process of a barrow of 100 or of 1 and zinc chardons. For example, the processing of the processing of the processing and processing the processin





Figure 1: BTA and LeA quantitation. A) 12% SDS-PAGE ged with BSA standards equivalent to 20, 15, 10, 08, 04, and 0.2 gg (lanes 2.7) was run in order to estimate the quantity of BTA and LeA. Three different amounts of BTA (lanes 9-11) and LEA (lanes 12-14) were sheed. After running the ged was scanned and analytered using a BisNatd G800 Densimeter. B) Shadhard curve obtained through plotting the density of the larow quantities of BSA. Or 10 µg vial of BTA was determined to contain 12.5 % -1.2 µg and one lor using 12.4 we determined to contain 12.3 % -04 µg.





Figure 3: Specific activity of RT. and LA using theorympic accept. Endoppoints on narring with SNAVAdd #530 were conducted in 20 and HEPSER (1814, 00.1 300 ACCRL) L356 MDT (140 and 0.3. WEPEN 20, 3. doayse with SNAVAdd #530 were conducted in 20 and HEPSE, 1814, 0.0.1 300 ACCRL) and 0.3. WEPEN 20, 0.3. The ACCR and 0.3. WEPEN 20, WEPEN



Figure 4: Transine d/ZaCi, in SNAPhile<sup>4</sup> massy. The SNAPhile<sup>4</sup> arrays were run ming SNA L(A) and § Jul SNAPhile<sup>4</sup> #SNA Were A) 50 mM HEPS, 816 8, 01 50 mM HEPS, 816 8-0. 059 MV HERD 2, 304 C) 50 mM HEPS, 816 8-0. 059 MV and 40 of reaction starting at 4 mM and intrang down to 1004 mM in 1: 84d dhiness. The machine were monitored continuously for one hor measuring BV over the starting at 4 mM and intrang down to 1004 mM in 1: 84d dhiness. The machine were monitored continuously for one hor measuring BV over history to the starting at 4 mM and intransition. Addition of the total of the other of the starting down of the first contact of the other oth



SSAPader 521 @ 3 µA
31 µM
(4.65 ng/ml)
25 µM
(1.25 ng/ml)

SSAPader 521@ 20 µA
No Data
25 µM
(1.25 ng/ml)

section of BTA and LeA. The limit of detection is defined as the minimum concentration of enzyme that can be measured by measured the section of users and the section of the section



Figure 6: Hydrolysie of VAMPick<sup>6</sup>, A) 8 μAV VAMPick<sup>6</sup> was claved by a series of concentrations of Lefl as indicated on the right. Digestions was performed in 50 mM HIRES, pH 6.2, containing 0.059 WTIEEN-201 at 37C. The lowest concentration, 0.318 M LeB, is easily detected in 160 minutes. B) Hydrolysis by LeB of either 8 μM substrate (red) or 20 μM substrate (blue) shows a linear response to ensyme concentration.



Figure 7: Addition of nite to the hydrolysis of VAMPile<sup>4</sup> and optimal pH. A) The charage of 8 pM VAMPile<sup>4</sup> by 20 AbLEs use performed in 50 mM (FIES) and 170, 47 × 40.8 A 2 containing 0.05% WTEXPS 201 at 740 percer Colsins. For all physics except 8, 2 ab best catalysis is seen without the addition of zinc. B) The hydrolysis of 8 pM VAMPile<sup>4</sup> by 20 AbLE as use toxed as a function of pH. The charage was performed in 50 nM MHS at PL 50, 57, and 53 containing 0.05% WTEXPS 201 at 74 he charage at pH 62, 43 5 6 6, 68, 71, 20 and 74 was performed in 50 nM MHS at PL 50, 57, and 53 containing 0.05% WTEXPS 201 at 74 he charage at pH 62, 43 5 6, 66, 87, 70, 24 and 74 was performed in 50 nM MHS at PL 50, 57, and 53 containing 0.05% WTEXPS 201 at 74 percent physics of 2 and 63, 45 6, 66, 87, 70, 70 and 74 was performed in 50 nM MHS at PL 50, 57, and 53 containing 0.05% WTEXPS 201 at 74 percent physics of 2 and 63, 45 6, 66, 87, 70, 70 and 74 was performed in 50 nM MHS at PL 50, 57, and 53 containing 0.05% WTEXPS 201 at 74 percent physics of 2 and 63, 45 6, 66, 87, 70 and 74 was performed in 50 nM MHS at PL 50, 57, and 53 containing 0.05% WTEXPS 201 at PL 50, 500 mL 50,

## **Conclusions**

### Specific Activity

The buildium neurotech type A fligt chain (LeA) has higher specific activity haved on units per namele enzyme, using babt SNAPidde'ubstratege when compared to the fling the survoirds if there is also a large fling fliences in activity have comparing the two FRAT substrates. SNAPidde\*2530 is a much hetter substrate than SNAPidde\*2531 is uben to Sing with either forms of the enzyme. We hypothesize that the large FRET pair on substrate 5521 may limit its access to the active site of the enzyme resulting in a discress in the number of annucle scales.

The specific activity of both LcA and BTA using SNAPtide<sup>®</sup> #520 appears to increase as enzyme concentration is decreased. We hypothesize that there may be some substrate inhibition occurring during these reactions. A portion of the cheaved substrate may interfere with further cleavage of the substrate.

### Addition of Zinc

20 mM HEPES buffer containing 0.05% TWEEN or 1 mg/ml BSA appears to be much more active than 20 mM HEPES buffer alone. We hypothesize that the TWEEN and BSA may help disperse non-specifically bound enzyme or substrate from the walls of the microfiler well. In so doing, more enzyme and substrate are free to interact.

The addition of high levels of aine chloride to the enzyme reactions are inhibitory. When using 20 mM HEPES with 0.05% TWEEN, no level of aine was breaficial. When the buffer system included 1 aging ISAs, how levels of aine chloride did improve the ultimate RFT achieved. Working with IRSAs in the buffer is somewhat problematic for the fact that bubbles are easily increduced in the week supen mixing and con affect the realing of the Bowerscence.

#### Limit of Detection (LOD)

The final of detection for bottlema warmstoric to pper 1 (algo chain is approximately) 2 pM ming there SNUM\*\* chainesses that are advector concentrations in a new rate of momental fits mass in the field of chain is adjusted of the filling bottlema mean chain (algo chain) is a specific of the fit of the fit

Our FRET substrates are a highly sensitive tool for detecting the activity of the botalinum neurotoxin. The sensitivity make it very useful in the search for small molecule inhibitors of the protease activity of the toxin.

#### VAMPtide®

VAMPide<sup>18</sup> is a new FRET substrate designed for testing activity of the B serotype of botalimum neurostoxia. It has been shown to be a good substrate for botalimum neurostoxia type B ight chain. The optimal pH is between 6.2 and 6.3 for the assay. The addition of ninc shoride to the assay was inhibitory, as was sense with LA. The rate of clearge of AVAPidie<sup>16</sup> is linear with respect to the occuentration of L4B.

Figure 5: Limit of detection of BTA and LoA. The limit of detection is defined as the minimum concentration of enzyme that can be measured with 99% confidence that the concentration is greater than acro under our testing conditions. A) The top graphs are representative calibration curves generated by testing BTA with SXNAHe #2014 at SM and when testing LAA with SXNAHe# 4520 at 2014. B) The limit of detection in this assay for LAA is among D42 pM. This value is true for both florongenic substrates and when using excess substrate. When compared to BTA the LOA D4 LoA is about 2014 both per or an and arcale.