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Identification of the synaptic vesicle glycoprotein 2 receptor binding site in botulinum neurotoxin A



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ABSTRACT

Botulinum neurotoxins (BoNTs) inhibit neurotransmitter release by hydrolysing SNARE proteins. The most important serotype BoNT/A employs the synaptic vesicle glycoprotein 2 (SV2) isoforms A-C as neuronal receptors. Here, we identified their binding site by blocking SV2 interaction using monoclonal antibodies with characterised epitopes within the cell binding domain (H_c). The site is located on the backside of the conserved ganglioside binding pocket at the interface of the H_{cc} and H_{cN} subdomains. The dimension of the binding pocket was characterised in detail by site directed mutagenesis allowing the development of potent inhibitors as well as modifying receptor binding properties.

Structured summary of protein interactions: H_cA binds to **rSV2A** by pull down (View interaction) **BONT/A** binds to **rSV2C** by pull down (1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11) H_cA physically interacts with **SV2A** and **SV2B** by anti tag coimmunoprecipitation (View interaction) H_cE physically interacts with **SV2A** and **SV2B** by anti tag coimmunoprecipitation (View interaction) H_cF physically interacts with **SV2A** and **SV2B** by anti tag coimmunoprecipitation (View interaction)

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1. Introduction

The fatal disease botulism, a flaccid paralysis of the muscles, is caused by blockade of the acetylcholine release at the neuromuscular junction due to intoxication with botulinum neurotoxins (BoNTs) even at minute quantities. On the other hand, of the seven major serotypes A-G the serotype BoNT/A constitutes a highly effective, widespread medicine to treat diseases like cervical dystonia, blepharospasm, hyperhidrosis and overactive bladder

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syndrome. All BoNTs are composed of four functional domains which play individual roles in their intoxication mechanism. The N-terminal 50 kDa light chain (LC) proteolyses specifically a member of the soluble N-ethlymaleimide sensitive attachment protein receptors family of proteins which are part of the vesicular fusion machinery [1,2]. The C-terminal 100 kDa heavy chain (HC) consists of three domains. Juxtaposed to LC is the 50 kDa α -helical translocation domain, the amino-terminal half of HC (H_N), and the remaining 50 kDa H_C-fragment comprises the H_{CN} domain which might trigger the translocation step [3] and the H_{CC} domain which harbours the binding sites for gangliosides and in case of BoNT/B, DC, and G for the synaptic vesicle (SV) protein receptor synaptotagmin [4].

BoNTs are taken up by neurons via a double receptor mechanism [5–7]. First, neuron specific abundant polysialo gangliosides like GT1b, GD1a and GD1b enrich BoNTs on the plasma membrane to facilitate contact with the intravesicular parts of the transiently occurring SV protein receptors [8]. E.g., BoNT/B, DC and G bind the membrane neighbouring 15–17 amino acids of the intraluminal

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Abbreviations: BoNT, botulinum neurotoxin; GBS, ganglioside binding site; GST, glutathion-S-transferase; HC, heavy chain; H_C, carboxyl-terminal half of HC; H_CA, H_CE, H_CF, H_CG, H_CX, H_C fragment of BoNT serotype A, E, F, G or A-G respectively; H_N, amino-terminal half of HC; H6F3H_CXS, N-terminal 6× His tag, 3× Flag tag, H_C fragment of serotype A-G, C-terminal Streptag; LC, light chain; mAb, monoclonal antibody; MPN assay, mice phrenic nerve hemidiaphragm assay; scFv, single-chain variable fragments; rSV2A, B, C, rat synaptic vesicle glycoprotein 2 isoform A, B, or C; Syt, synaptotagmin; CD, circular dichroism

domain of the synaptic vesicle type I membrane proteins synaptotagmin (Syt)-I and Syt-II in the distal tip of their H_C-fragment (Syt site) in the direct neighbourhood of the conserved ganglioside binding pocket in different arrangements [6,9–16].

Later, the large intravesicular domain 4 (LD4) of the synaptic vesicle glycoprotein 2 (SV2) isoforms A–C was discovered as receptor domains for BoNT/A [17,18], the unglycosylated, isolated LD4 of SV2C exhibiting the highest affinity to BoNT/A H_c. However, since all three SV2 isoforms are expressed in α -motoneurons [17], it is still not clear which SV2 isoform is most relevant for the physiological uptake of BoNT/A.

In contrast to BoNT/A, functional studies revealed that BoNT/E only interacts with SV2A and SV2B, but not with SV2C [19] while the biological activity of BoNT/D primarily requires the presence of SV2B and to a lesser extent of SV2A and C [20]. However, a direct protein–protein interaction between SV2 and BoNT/D remains to be shown like for BoNT/F which co-precipitated SV2A-C from rat brain synaptic vesicle lysates [21,22]. Altogether, BoNT/A, D, E and F utilise seemingly diverse SV2-binding mechanisms.

Only recently, the first SV2 binding pocket could be mapped in BoNT/E to an extended surface area located on the back of the conserved ganglioside binding site comprising the H_{CN} and H_{CC} interface, part of the corresponding Syt site in BoNT/B and the region in between (ODA1 site), the latter two also constituting a partial epitope of the monoclonal antibody (mAb) 4E13 which neutralises BoNT/E activity [23]. The SV2A binding site in BoNT/E seems to represent the prototypic region for all other BoNTs that exploit SV2 as a cell surface receptor. In accordance, a co-crystal structure allocated the binding site of human SV2C in BoNT/A to the H_{CN} and H_{CC} interface adjacent to the SV2A binding site in BoNT/E [24].

Here, we identified the SV2 binding site at the interface of H_{CC} and H_{CN} domains of BoNT/A employing neutralising mAbs with defined epitopes used as therapeutics [25]. Site-directed mutagenesis characterised the SV2 binding pocket in detail and exhibited the single mutation G1292R decreasing the biological activity of BoNT/A ~300-fold. Employment of ganglioside deficient mice and BoNT/A mutants comprising deactivated receptor binding sites proved that the double receptor model also accounts for BoNT/A.

2. Materials and methods

Anti-BoNT/A mAb CR2 and scFvs RAZ1, S25 and CR2 were generated as described previously [25–29]. Fig. 1B was prepared using the program Discovery Studio Visualizer 2.5 (Accelrys Software Inc.) based on 2NYY.pdb [28].

2.1. Plasmid constructions and production of proteins

Rat GST-rSV2C 454–579 encoded by pGEXrSV2C 454–579 and rat GST-rSV2A 468–594 encoded by pGEXrSV2A 468–594 were produced as described previously [18]. The plasmid pH6tBoNTA1 encoding BoNT/A1 fused to an N-terminal His6tag was generated by modifying pBoNTAS [30]. The plasmids pH_cAS, pH6F3H_cES and pH6F3H_cGS were described previously [15,23,30], pH6F3H_cES serving as parental vector for pH6F3H_cAS and pH6F3H_cFS which were generated by ligation of DNA sequences encoding H_cA (aa 871–1296) from pS3H_cAH6 and H_cF (aa 866–1277) from pS3H_c-FH6 [22]. All pH6tBoNTA1 and pH6F3H_cAS mutants were prepared using the GeneTailor method (Invitrogen GmbH, Karlsruhe, Germany) employing suitable primers and pH6tBoNTA1 and pH6F3H_c-AS, respectively, as template DNA. Nucleotide sequences of all mutants were verified by DNA sequencing.

Wild-type and mutated recombinant H6F3H_CXS (= N-terminal $6 \times$ His tag, $3 \times$ Flag tag, H_C fragment of serotype A-G, C-terminal Streptag), H_CAS and full-length neurotoxins, the latter under

biosafety level 2 containment (project number GAA A/Z 40654/3/ 123), were produced utilising the Escherichia coli strain M15pREP4 (Qiagen, Germany) during 16 h of induction at 22 °C in the presence of 0.2 mM IPTG, and were purified on Co²⁺-Talon matrix (Takara Bio Europe S.A.S., France). Full-length neurotoxins were eluted using 50 mM Tris-HCl, pH 8.0, 150 mM NaCl, 250 mM imidazole, subjected to gel filtration (Superdex-200 16/60 column, GE Healthcare, Germany) in 100 mM Tris-HCl, pH 8.0, 150 mM NaCl, frozen in liquid nitrogen and kept at -70 °C. H6F3HcXS proteins were eluted using 100 mM Tris-HCl pH 8.0, 150 mM NaCl, 100 mM Imidazole and further purified on StrepTactin-Sepharose beads (IBA GmbH, Germany). Proteins were eluted by 10 mM desthiobiotin in 100 mM Tris-HCl pH 8.0, 150 mM NaCl, frozen in liquid nitrogen and kept at -70 °C. For circular dichroism (CD) analysis desired volume of protein was dialysed against $1 \times PBS$ pH 7.4.

Protein concentrations were determined subsequent to SDS– PAGE and Coomassie blue staining by using a LAS-3000 imaging system (Fuji Photo Film), the AIDA 3.51 program, and various known concentrations of BSA as reference.

2.2. GST-pull down assays

For competition experiments H_cAS (100 pmol) or H6tBoNTA (50 pmol) were preincubated with mAb (400 pmol) for 30 min at RT in 187.5 μ l in 100 mM Tris–HCl, pH 8.0, 150 mM NaCl.

Glutathion-S-transferase (GST, 150/75 pmol), GST-rSV2A 468– 594 (150 pmol) and GST-rSV2C 454–579 (75 pmol) immobilised to 10 μ l of glutathione-Sepharose-4B matrix (GE healthcare, Germany) were incubated with H_cAS (100 pmol), H6tBoNTA wild-type (with or without preincubated mAbs) or mutants (50 pmol each) in a total volume of 200 μ l 100 mM Tris–HCl, pH 8.0, 150 mM NaCl, supplemented with 0.5% Triton X-100 (Tris/NaCl/Triton) buffer for 2 h at 4 °C. Beads were collected by centrifugation and washed two times each with 200 μ l Tris/NaCl/Triton buffer. Washed pellet fractions were denatured in SDS sample buffer for 20 min at 37 °C and analysed by SDS–PAGE. Proteins were detected by Coomassie blue staining and subsequently quantified by densitometry.

2.3. Mouse phrenic nerve hemidiaphragm (MPN) assay

The mice phrenic nerve hemidiaphragm assay (MPN assay) was performed as described previously employing 20-30 g NMRI mice (Janvier SA, France) [31] or complex polysialo ganglioside deficient C57BL/6 mice lacking the genes B4gaInt1 and/or St8sia1 [22]. The phrenic nerve was continuously stimulated at 5-25 mA with a frequency of 1 Hz and with a 0.1 ms pulse duration. Isometric contractions were transformed using a force transducer and recorded with VitroDat Online software (FMI GmbH, Germany). The time required to decrease the amplitude to 50% of the starting value (paralytic half-time) was determined. To allow comparison of the altered neurotoxicity of mutants with H6tBoNTA wild-type, a power function (y(H6tBoNTA; 10, 30, 80 pM) = $139.6x^{-0.1957}$ $R^2 = 0.9991$) was fitted to a concentration-response-curve consisting of three concentrations determined minimum in triplicates. Resulting paralytic half-times of the H6tBoNTA mutants were converted to concentrations of the wild-type employing the above power functions and finally expressed as relative neurotoxicity (Figs. 3 and 5).

2.4. Co-immunoprecipitation

All centrifugation and incubation steps were carried out at 4 °C, except where otherwise stated. The preparation of rat brain synaptosomes was performed as previously described [31].



Fig. 1. (A) Binding of full-length BoNT/A to GST-rSV2C 454–579 is blocked by monoclonal antibody IgG CR2 and its corresponding scFv. H6tBoNTA (50 pmol) serving as prey was preincubated with mAb (400 pmol) for 30 min at RT ("input") in 200 µl 100 mM Tris–HCl, pH 8.0, 150 mM NaCl, 0.5% Triton X-100 and subsequently incubated for 2 h at 4 °C with the bait, immobilised GST-rSV2C 454–579 (75 pmol). Bound H6tBoNTA was detected by 12.5% SDS–PAGE analysis and Coomassie blue staining. (B) Complex crystal structure of BoNT/A H_C (H_{CC}, light gray surface; H_{CN}, dark grey surface) and the variable part of CR1 Fab (orange ribbon; modified from 2NYY.pdb). Residues in the analogous Syt site/RA21 epitope are depicted in green, those of the ODA1 site in light blue or dark blue (effect on SV2C binding). All positions analysed in the CR2 site are coloured in orange, those with clear effect on SV2C binding in red.

Synaptosomes were solubilised in lysis buffer (20 mM Tris–HCl, pH 7.4, 80 mM NaCl, 0.5% Triton X-100) for 1 h, subsequently centrifuged at 21,000×g for 10 min and the supernatant was transferred into a fresh tube. Co-immunoprecipitation of native SV2A and B was done in lysis buffer supplemented with 0.1% BSA in a total volume of 200 μ l employing 10 μ l Protein G agarose beads (Amersham Biosciences), 6 μ g of anti-Flag M2 monoclonal antibody (Sigma–Aldrich), 12 μ g of the indicated BoNT H_c-fragment and

50 μ g of synaptosomal proteins in the presence of 125 μ g mixed bovine brain gangliosides for 3 h. Thereafter, beads were collected by centrifugation at 2000 \times g and washed three times using lysis buffer. Washed pellet fractions were incubated in Laemmli sample buffer for 20 min at 37 °C and analysed by SDS–PAGE and Western blotting (rabbit polyclonal anti-SV2A & anti-SV2B, Synaptic Systems, Germany; goat anti-rabbit, Pierce; StrepMAP Classic-HRP, IBA GmbH, Germany).



Fig. 2. Binding of H6tBoNTA single mutants (50 pmol) for 2 h at 4 °C to immobilised GST-rSV2C 454–579 (75 pmol) in 200 µl 100 mM Tris–HCl, pH 8.0, 150 mM NaCl, 0.5% Triton X-100. Bound H6tBoNTA mutants were detected by 10% SDS–PAGE analysis and Coomassie blue staining (wt *n* = 15, mutants *n* = 3–5, ±SD), quantified by densitometry and compared versus wild-type binding.



Fig. 3. The impact of mutations on the neurotoxicity of BoNT/A was analysed employing MPN preparations. The paralytic halftimes were determined and converted to the corresponding concentrations of wild-type BoNT/A using a concentration–response curve. The resulting toxicities were finally expressed relative to wild-type BoNT/A (*n* = 3–5, ±SD).

3. Results and discussion

The H_C-fragment of BoNT/A displays the highest affinity to the luminal domain 4 (residues 454–579) of the isoform SV2C independent of its glycosylation status followed by SV2A and SV2B [17,18]. This direct protein–protein interaction is clearly demonstrated in a GST pull down assay employing full-length recombinant H6tBoNTA and rat GST-rSV2C 454–579 (Fig. 1A). The weaker binding of H_CA to recombinant rat GST-rSV2A 468–594 is also shown (Fig. S1) whereas binding to non-glycosylated rSV2B is hardly detectable. Well characterised neutralising monoclonal

antibodies directed against different epitopes within the H_c -fragment of BoNT/A [25–29] were screened for interference with the SV2C interaction. To avoid putative cross-linking of two BoNT/A molecules by an intact IgG which could impair SV2 binding, we initially pre-incubated H6tBoNTA with the single-chain variable fragments (scFv) S25, RAZ1 and CR2 in 8-fold excess. scFv S25 is directed against residues 1254–56 forming a part of the rim of the single ganglioside binding site (GBS) [30]; RAZ1 is an affinity matured mAb based on 3D12 directed against residues 1129–31 comprising the region analogous to the Syt binding site in BoNT/ B and G [6,14]. CR2 and the closely related CR1 are engineered



Fig. 4. Co-immunoprecipitation of SV2A and B by selected H_cA mutants was performed by incubating 12 µg of 3 × Flag-tagged wild-type and mutant H6F3H_cAS, H6F3H_cES, H6F3H_cFS and H6F3H_cCS with 6 µg anti-Flag-tag M2 antibody and Triton X-100 solubilised rat brain synaptosomal membranes. Bound SV2A and B were detected by Western blotting, amount of precipitated H6F3H_cX was verified by HRP conjugated StrepMAB classic. MWM denotes molecular weight marker.



Fig. 5. The neurotoxicity of wild-type and BoNT/A mutants either with inactivated GBS (W1266L), inactivated SV2 binding site (R1156M, G1292R) or both was determined employing wild-type and ganglioside-deficient tissue for the MPN assay ($n = 7-16, \pm$ SD).

mAbs derived from C25 which binds predominantly to residues 918–20, F953 and 1062–64 at the interface of the H_{CC} and H_{CN} domain [28,29] (Fig. 1B). Whereas scFv S25 did not and RAZ1 only marginally altered the interaction of GST-rSV2C with H6tBoNTA, the scFv CR2 eliminated the precipitation of H6tBoNTA by GST-rSV2C (Fig. 1A). Analogously, the 48-fold excess of IgG CR2 terminated binding of H6tBoNTA to GST-rSV2C, which explains

its potency in *ex vivo* MPN assay as well as in *in vivo* mice toxin neutralisation assays [26]. Hence, a main interaction site of SV2C with BoNT/A resides in the epitope region of CR2. Analogously, CR2 was able to inhibit the precipitation of H_cA by GST-rSV2A LD4 (468–594) indicating that both SV2 isoforms interact with the same surface area of BoNT/A (Fig. S1). Previously, the CR2 epitope was mapped in detail by two crystal structure complexes consisting of BoNT/A and Fabs of CR1 or AR2, respectively, closely related mAbs recognising identical epitopes [28] (Fig. 1B). Based on this structural data site directed mutagenesis was performed in the CR2 site, the Syt/RAZ1 site as well as in the region connecting both surface areas called ODA1 site according to Mahrhold et al. [23].

The full-length recombinant H6tBoNTA mutants were screened in pull down assays employing rat GST-rSV2C 454–579. Charge reversal by R1131E in the RAZ1 site and K1163E in the ODA1 site decreased precipitation of H6tBoNTA, even more pronounced for mutation N1188A in the ODA1 site (Fig. 2). Furthermore, 15 mutations in the CR2 site were generated. Here, the H6tBoNTA single mutants N905Y, T1063P, H1064G, D1108R, R1156M, V1287E and G1292R displayed a 50% or greater decrease of binding to GSTrSV2C which is in good agreement with binding assay data obtained for the CR2-related humanised C25 [29] and the crystallised complexes BoNT/A-CR1 or AR2 which identified i.a. T1063 and H1064 as key residues in their epitopes [28]. With the exception of D1124R also the remaining seven mutants showed at least moderately reduced affinity to GST-rSV2C (Fig. 2).

To verify whether the reduced affinity of the BoNT/A mutants to recombinant rSV2C LD4 also translates into reduced biological activity all 21 mutants were tested in the MPN assay. Correspondingly, mutants H6tBoNTA R1131E and K1260E in the RAZ1 site and N1188A within the ODA1 site showed diminished potency (Fig. 3). Also mutations N905Y, F917A, T1063P, H1064G, D1108R, P1110W, D1124R, R1156M, V1287E and D1289G/R in the CR2 site caused a decrease of more than 50% in neurotoxicity. The single mutant G1292R even displayed a 300-fold reduction in potency. Thus the MPN data largely confirms the previous in vitro binding data obtained with isolated rat GST-rSV2C 454-579, also because the LD4 of mouse and rat SV2C differ only in three conservative amino acid exchanges. Noteworthy, the isoforms SV2A and B are also expressed in motoneurons [17], although neither their differential expression levels nor their affinity to BoNT/A have been quantified. The amino acid sequence identity between the LD4 of rSV2A, rSV2B and rSV2C is only 48% and 44%, respectively. Hence, differences between the GST-rSV2C pull down and the MPN assay as observed for mutants P1110W, D1124R and D1289G/R could be explained by subtle differences in the mode of interaction of the three SV2 isoforms. In conclusion, the main SV2C binding site is allocated in the CR2 epitope site at the interface of H_{CC} and H_{CN} domains on the backside of the GBS extending towards the Syt/RAZ1 binding site. Our finding is in good agreement with a recently published co-crystal structure of human SV2C residues 456–579 bound to the BoNT/A H_C-fragment where one β -strand of the SV2C quadrilateral β -helix interacts with the convex β -sheet (1140-56) at the BoNT/A H_{CC}-H_{CN} interface [24]. This β -sheet is adjacent to the CR2 epitope in H_{CN}-H_{CC}. Here, T1145/ 46 and R1156 dominate the few proposed side chain-side chain interactions. Analogously, our mutant R1156M showed the lowest affinity to GST-rSV2C. Secondly, the strong effects observed for mutation of G1292 is in accordance with data of Benoit et al. showing impaired hSV2C interaction upon mutation of the neighbouring R1294 to alanine [24]. Furthermore, comparison with the SV2Abinding site in BoNT/E shows that mutation of Y879, E1246 and K1084 in BoNT/E, like the corresponding residues N905, V1287 and D1108 in BoNT/A for SV2C, interferes with SV2A-binding of BoNT/E. This illustrates that the SV2 binding site of BoNT/A covers the homologous region of BoNT/E [23].

To ensure that the single mutations on the H_C-fragment surface did not impair its structural integrity and thereby caused the loss in SV2C binding, the mutations displaying the most severe effects (R1156M, V1287E and G1292R) were transferred into H_C-fragments fused to an N-terminal 3 × FLAG tag. The thermal stability of the H_C-fragment mutants was virtually identical to the wild-type (51 ± 1 °C; Fig. S2) indicating that the secondary structure was not altered due to the single amino acid exchanges.

To further deepen the conclusion obtained for the interaction site of SV2 in BoNT/A, co-immunoprecipitation experiments were conducted employing $3 \times$ FLAG tagged BoNT/A H_C-fragment mutants (H6F3H_cAS; Fig. 4). Since the tissue from motoneurons as physiological target cannot be accessed in sufficient quantities, rat brain synaptosomes were generated and solubilised in Triton X-100. However, whereas SV2A is ubiquitously expressed in virtually all synapses and SV2B was also found in a wide variety of synapses throughout the brain, SV2C was observed only in few brain areas like the pallidum, the substantia nigra, the midbrain, the brainstem and the olfactory bulb, but was undetectable in the cerebral cortex and the hippocampus, and found at low levels in the cerebellar cortex [32]. Hence, SV2C constitutes only a minor fraction in synaptosomal lysates and H_cA wild-type co-precipitated only minor amounts. In contrast, SV2A and SV2B were satisfactorily precipitated by H_cA wild-type (Fig. 4). H_cA mutants R1156M and V1287E showed similar affinity to SV2A and SV2B as wild-type H_cA indicating subtle differences between the mode of interaction of SV2A/B and of SV2C with BoNT/A. R1156 is suggested to interact with F563 of human SV2C by cation $-\pi$ -stacking interaction [24]. However, mouse and rat SV2C comprise L563 which does not allow this cation– π -stacking, but possibly R1156 establishes a salt bridge to the neighbouring E543 in rat SV2C. However, the corresponding residues Y557 in rSV2A and Y500 in SV2B would form a weaker H-bond to R1156. Consequently, mutant R1156M does only moderately interfere with SV2A and B binding. The lack of effect for mutant V1287E cannot be deduced from the H_cA-hSV2C complex structure. In contrast, mutant H_cA G1292R did neither co-precipitate SV2A nor SV2B which confirms the conclusion that also SV2A and SV2B interact at the similar region on the H_cA surface. Hereby, the strong effect of mutation G1292R on SV2A-C interaction also explains the drastic decrease of biological activity of H6tBoNTA G1292R in the MPN assay compared to all other mutants in the CR2 site.

The single BoNT/A mutant G1292R with eliminated SV2 protein receptor recognition, the mutant W1266L lacking ganglioside binding [30] as well as ganglioside deficient mice [33] provided powerful tools to prove the dual receptor hypothesis for BoNT/A analogously to BoNT/B and G [6]. Like previously shown [22], the sensitivity of tissue expressing GM3 only was decreased by \sim 600-fold for wild-type H6tBoNTA (Fig. 5) which corresponds to the decrease in sensitivity of wild-type mice hemidiaphragm tissue towards H6tBoNTA W1266L bearing a deactivated GBS [30] (Fig. 5). Whereas the neurotoxicity of the mutant W1266L at GM3-only tissue remained roughly the same, the H6tBoNTA G1292R with inactivated SV2 binding site (= 300-fold reduced neurotoxicity at wildtype tissue) showed further reduced neurotoxicity at gangliosidedeficient tissue indicating the importance of both receptor interactions (Fig. 5). Finally, the double mutant comprising W1266L and G1292R turning both receptor sites inactive displayed a 250,000fold reduced biological activity in the MPN assay (Fig. 5). In comparison, the individual mutations each decreased the potency by \sim 200- and 300-fold demonstrating the synergy of both binding sites for the intoxication mechanism.

In conclusion, exchange of two critical amino acids in the GBS and the newly identified SV2 binding site generates a virtually atoxic BoNT/A which adds another piece of evidence for the dual receptor mechanism by which botulinum neurotoxins specifically enter nerve cells [4,5]. Furthermore, the knowledge of both recep-

tor binding sites now allows the design of bivalent inhibitor molecules blocking the uptake of BoNT/A in case of its deliberate or accidental release. Also the retargeting of BoNT/A to e.g. non-neuronal cells to treat diseases caused by overactive secretion mechanisms which has so far been achieved by replacing the whole H_{C} fragment [34] could now be accomplished with minimal alterations in the SV2 binding site. And in combination with three critical substitutions in the LC interfering with the catalytic mechanism [35–37] such a quintuple mutant represents an ideal atoxic antigen to generate vaccines against physiologically folded BoNT/A.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.febslet. 2014.02.034.

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