**Cell Chemical Biology**

**Differential effects of modified batrachotoxins on voltage-gated sodium channel fast and slow inactivation**

*Graphical abstract*

*Highlights*

- BTX and C20-ester derivatives modulate gating of sodium channel subtypes

- BTX hyperpolarizes voltage of activation and inhibits both fast and slow inactivation

- A BTX C20-ester differentiates fast and slow inactivation, only blocking the former

*Authors*


*Correspondence*

jdubois@stanford.edu (J.D.), daniel.minor@ucsf.edu (D.L.M.)

*In brief*

BTX, an acute poison, acts as a potent allosteric regulator of NaVs. Structure-function analysis of BTX and three BTX C20-ester derivatives reveals unexpected differences in activity. One compound, BTX-yne, blocks fast but not slow inactivation of ion conduction and is thus distinguished from the others.
Differential effects of modified batrachotoxins on voltage-gated sodium channel fast and slow inactivation

Tim M.G. MacKenzie,1,7,8 Fayal Abderemane-Ali,2,7 Catherine E. Garrison,1,7 Daniel L. Minor, Jr.,2,3,4,5,6,* and J. Du Bois1,9,*

1Department of Chemistry, Stanford University, 337 Campus Drive, Stanford, CA 94305, USA
2Cardiovascular Research Institute, University of California, San Francisco, Box 3122, 555 Mission Bay Boulevard South, Rm. 452Z, San Francisco, CA 94158-9001, USA
3Departments of Biochemistry and Biophysics, and Cellular and Molecular Pharmacology, University of California, San Francisco, CA 94158-9001, USA
4California Institute for Quantitative Biomedical Research, University of California, San Francisco, CA 94158-9001, USA
5Kavli Institute for Fundamental Neuroscience, University of California, San Francisco, CA 94158-9001, USA
6Molecular Biophysics and Integrated Bio-imaging Division, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, USA
7These authors contributed equally
8Present address: Department of Genetics, Stanford University, 3165 Porter Drive, Palo Alto, CA 94304, USA
9Lead contact
*Correspondence: jdubois@stanford.edu (J.D.), daniel.minor@ucsf.edu (D.L.M.)
https://doi.org/10.1016/j.chembiol.2021.12.003

SUMMARY

Voltage-gated sodium channels (NaVs) are targets for a number of acute poisons. Many of these agents act as allosteric modulators of channel activity and serve as powerful chemical tools for understanding channel function. Herein, we detail studies with batrachotoxin (BTX), a potent steroidal amine, and three ester derivatives prepared through de novo synthesis against recombinant NaV subtypes (rNaV1.4 and hNaV1.5). Two of these compounds, BTX-B and BTX-CHx, are functionally equivalent to BTX, hyperpolarizing channel activation and blocking both fast and slow inactivation. BTX-yne—a C20-n-heptynoate ester—is a conspicuous outlier, eliminating fast but not slow inactivation. This property differentiates BTX-yne among other NaV modulators as a unique reagent that separates inactivation processes. These findings are supported by functional studies with bacterial NaVs (BacNaVs) that lack a fast inactivation gate. The availability of BTX-yne should advance future efforts aimed at understanding NaV gating mechanisms and designing allosteric regulators of NaV activity.

INTRODUCTION

Voltage-gated sodium channels (NaVs) are central to physiological function as requisite protein complexes that drive the initiation and propagation of action potentials in electrically excitable cells (Hille, 2001; Peters and Ruben, 2014; Chahine, 2018). NaV malfunction underlies a number of human pathologies, and, unsurprisingly, acute disruption of NaV activity can be fatal (Ashcroft, 1999; George, 2005; Fiske, 2006; Catterall, 2008; Lampert, 2010). A large and structurally disparate collection of poisons occur in nature that target NaVs (Stevens et al., 2011; Kalia, 2015; Ahern, 2016; Deuis, 2017; Lukowski and Narayan, 2019). The lethality of these agents notwithstanding, peptide and small molecule toxins have proven invaluable as chemical reagents for studies of NaV structure and dynamics. Among such molecules, one natural product—batrachotoxin (BTX)—is conspicuous. BTX is a steroidal amine derivative commonly associated with poison dart frogs, but also found in species of bird and beetle (Khodorov, 1988; Brown, 1988; Dumbacher et al., 2000, 2004). This toxin is unique among all other NaV modulators in that it affects every measurable aspect of NaV function including activation threshold, inactivation, single-channel conductance, and ion selectivity (Catterall et al., 1981; Brown, 1988; Linford et al., 1998; Bosmans, 2004). Because of its chemical complexity and lack of availability from natural sources, studies to dissect how the different structural elements of BTX contribute to altering channel function are limited (Brown et al., 1981; Khodorov et al., 1992; Garraffo and Spande, 2009; Yelin et al., 2019). De novo synthesis, however, has enabled access to (-)-BTX and modified forms thereof (Logan, 2016). In this report, experiments comparing the effects of BTX and three BTX C20-ester derivatives against skeletal muscle and cardiac NaVs reveal a singular agent, BTX-yne, that differentially influences fast and slow channel inactivation. This unique property distinguishes BTX-yne as a powerful tool for biophysical and pharmacological studies of NaVs. The availability of BTX-yne should advance efforts to understand the mechanisms...
underlying Na\textsubscript{v} gating and guide the development of allosteric modulators for treating Na\textsubscript{v}-related disorders.

**Background**

Eukaryotic Na\textsubscript{v}s are heteromeric protein complexes consisting of a pore forming \(\alpha\)-subunit and up to two auxiliary \(\beta\)-proteins (Catterall, 2012; Ahern, 2016; Chahine, 2018). The \(\alpha\)-subunit, of which there are nine different isoforms (Na\textsubscript{v}1.1–1.9), derives from a single polypeptide chain that clusters into four homologous repeats (DI–DIV), each consisting of six transmembrane \(\gamma\)-helices (S1–S6). Functionally, Na\textsubscript{v}s have two “gates” (activation and inactivation gates), both of which must be open in order for ion conduction to occur (Bezanilla, 2008; Catterall et al., 2017).

The simplest model to describe Na\textsubscript{v} gating posits three limiting conformations: 1) a non-conducting closed state that is primed to respond to membrane depolarization; 2) an open, conducting state; and 3) a non-conducting inactivated state (Patlak, 1991). Inactivated channels must transition back to a closed form before firing again. A sufficiently strong membrane depolarization induces the outward movement of the voltage-sensors, causing channels to open (activation). Opening is rapidly followed by inactivation, which occurs on the millisecond timescale (fast inactivation). Fast inactivation involves the movement of a hydrophobic IFM motif located on a loop connecting DIII–DIV, which allosterically effects pore closure (West, 1992; Pan, 2018). An alternative form of Na\textsubscript{v} inactivation (slow inactivation) takes place over longer time scales (seconds to minutes) and results from repetitive or sustained membrane depolarization (Vilim et al., 2001; Silva, 2014). Additionally, channels can directly inactivate from the closed state without ever entering the open state (closed-state inactivation) (Goldman, 1995; Armstrong, 2006). The steady-state distribution of closed, open, and inactivated channels at a resting membrane potential is intrinsic to each Na\textsubscript{v} \(\alpha\)-subunit. Natural toxins and other small molecules that bind the \(\alpha\)-subunit are generally biased toward a particular conformational state of the channel and can affect one or more properties such as ion conduction, channel activation, inactivation, and ion selectivity (Wang and Wang, 2003; Stevens et al., 2011; Kalia, 2015; Ahern, 2016; Deuis, 2017; Lukowski and Narayan, 2019). Studies with such agents have been instrumental in dissecting the complex dynamics of channel gating.

Among the most venerable Na\textsubscript{v} toxins is the steroidal derivate Batrachotoxinin A (BTX-A), which allosterically effects pore closure (West, 1992; Pan, 2018). An alternative form of Na\textsubscript{v} inactivation (slow inactivation) takes place over longer time scales (seconds to minutes) and results from repetitive or sustained membrane depolarization (Vilim et al., 2001; Silva, 2014). Additionally, channels can directly inactivate from the closed state without ever entering the open state (closed-state inactivation) (Goldman, 1995; Armstrong, 2006). The steady-state distribution of closed, open, and inactivated channels at a resting membrane potential is intrinsic to each Na\textsubscript{v} \(\alpha\)-subunit. Natural toxins and other small molecules that bind the \(\alpha\)-subunit are generally biased toward a particular conformational state of the channel and can affect one or more properties such as ion conduction, channel activation, inactivation, and ion selectivity (Wang and Wang, 2003; Stevens et al., 2011; Kalia, 2015; Ahern, 2016; Deuis, 2017; Lukowski and Narayan, 2019). Studies with such agents have been instrumental in dissecting the complex dynamics of channel gating.

Among the most venerable Na\textsubscript{v} toxins is the steroidal derivate, BTX, the active component in poison darts obtained from frog secretions by Colombian natives (Mark and Witkop, 1963; Daly, 1965; Tokuyama, 1968, Tokuyama et al., 1969). BTX is the prototypical member of the family of lipid soluble toxins that lodge in the inner pore of the channel (termed site II). These toxins are unique in simultaneously binding to the inner pore and disrupting channel gating. Notably, BTX binding affects every measurable aspect of Na\textsubscript{v} function: the activation threshold is shifted in the hyperpolarizing direction by 30–50 mV, both fast and slow inactivation are abolished, single-channel conductance is reduced, and ion selectivity is compromised (Linford et al., 1998; Bosmans, 2004; Wang et al., 2006). Efforts to understand BTX activity have relied on electrophysiology in combination with protein mutagenesis and structure-activity relationship (SAR) studies (Linford et al., 1998; Wang and Wang, 1998, 1999, 2017; Wang et al., 2000, Wang et al., 2001, Wang, 2007; Li et al., 2002; Du et al., 2011). Ligand docking studies with homology models of the Na\textsubscript{v} pore suggest specific contacts between toxin and channel but, to date, reveal little about toxin function. The studies detailed herein are enabled by our ability to access BTX through a multistep synthesis (Logan, 2016). This work has led to the identification of BTX-\textsubscript{yne}, a BTX derivative that alters Na\textsubscript{v} gating by inhibiting fast, but not slow, inactivation. This characteristic is distinct from BTX and two other C20-ester variants, all three of which block both fast and slow inactivation. To our knowledge, no other small molecule or peptide has been described that separates these two functional features of the channel.

**RESULTS**

**Synthesis of BTX derivatives**

Following our reported synthesis of batrachotoxinin A (BTX-A), we have optimized conditions for modifying the C20 alcohol in order to install different ester groups (Scheme 1). The ester moiety is essential for toxin activity, as BTX-A is > 1,000 times less potent than the parent compound (Albuquerque et al., 1971; Brown, 1988). For the purpose of this study, four compounds were prepared, BTX, BTX-B, BTX-\textsuperscript{2}Hx, and BTX-\textsuperscript{yne}, and evaluated against recombinant rat skeletal muscle sodium channels (rNa\textsubscript{v}1.4). The benzoate ester, BTX-B, has been reported previously and is generally regarded as a functional equivalent of BTX (Brown et al., 1981; Catterall et al., 1981); however, a direct quantitative comparison of BTX and BTX-B compounds has not been detailed. The other two derivatives, BTX-\textsuperscript{2}Hx and BTX-\textsuperscript{yne}, were designed to assess the importance of the aryl moiety for toxin activity against Na\textsubscript{v} (Tikhonov and Zhorov, 2005; Du et al., 2011). We note that BTX-\textsuperscript{yne} and BTX-\textsuperscript{2}Hx were selected over other ester groups, as these two derivatives are comparable to BTX and BTX-B in terms of lipophilicity based on similar cLogP values (BTX 3.22; BTX-B 4.12; BTX-\textsuperscript{yne} 3.58;
BTX derivatives differentially influence NaV inactivation

A signature effect of BTX on NaV activity is the elimination of both fast and slow inactivation (Figure 1A). To determine if C20-ester substitution of the toxin alters this behavior, we used a two-pulse protocol to measure steady-state inactivation (SSI) (Figure 2 inset) (Bendahhou et al., 1999). Channels that inactivate in the pre-pulse period are unavailable to open in the test pulse. A plot of normalized current versus voltage (I/I_0—V) for unmodified NaV1.4 channels (black circles) under this SSI protocol is shown in Figure 2C. Analogous SSI curves for toxin-treated NaVs display similar Boltzmann behavior, but a fraction of the total population of channels is non-inactivating; this percentage varies in a dose-dependent manner (Figures S3 and D2, Table S3). At saturating concentrations of BTX, BTX-B, and BTX-^2Hx, inactivation is completely abrogated (Figures 2A, S3, D2). EC_{50} values for the SSI response were determined by varying toxin concentration (Table 1). These values accord with voltage of activation EC_{50s} for each compound.

Comparison of SSI curves recorded at saturating concentration (10 μM) of each of the four toxin derivatives reveals an unexpected difference for BTX-yne (Figures 2A—2C, S3). In contrast to BTX, BTX-B, and BTX-^2Hx, which eliminate inactivation, BTX-yne-modified channels are able to inactivate partially. Interestingly, a plot of I/I_0 versus voltage appears as a U-shaped SSI curve (vide infra). The extent to which BTX-yne-treated channels
are inactivated can be estimated by applying a baseline correction to the SSI curve (Figure S4). From this analysis, 40%–50% of channel inactivation occurs at a saturating concentration of BTX-yne, as compared with ca. 100% and 0% for untreated and BTX-B-modified channels, respectively. The influence of BTX-yne on SSI is not unique to rNaV1.4, as hNaV1.5 channels treated with this compound (5 μM) are similarly altered (Figure S5).

Although all four toxin derivatives shift the activation voltage dependence to a comparable extent, BTX-yne stands alone as the only compound that does not completely inhibit channel inactivation (Figures 1C and 2C). Superposition of the voltage-dependent activation and inactivation curves shows that, in the presence of BTX-yne, inactivation initiates at potentials at which channels are closed (<−100 mV) and persists even when all channels are open (>−40 mV) (Figures S4E–S4F). This observation suggests that BTX-yne-modified channels undergo inactivation from both the closed and open states (i.e., closed-state and open-state inactivation, respectively).

Recording data shown in Figure 1B illustrate that BTX-yne blocks NaV1.4 fast inactivation, analogous to the other three agents (Figures S3 and D2). The question thus arises: can fast inactivation, the latter stemming from the large variable tail currents, which interfere with measurements in the test pulse. Cells were stepped from a holding potential of −120 to −70 mV for varying lengths of time, Δt. Following this conditioning pulse, the extent to which channels inactivate as a function of Δt was measured in a test pulse to 0 mV (Figure S6) (Cummins et al., 1998). BTX-B was employed as a negative control for these experiments, as channels bound to this compound remain conducting irrespective of the time period of the conditioning or test pulses. Using this stimulation protocol, cells treated with 10 μM BTX-yne display a clear exponential decay of peak current as a function of the conditioning pulse length. Fitting these data for BTX-yne to two exponentials gives a time constant, τ, for each process (τfast = 90.8 ± 3.8 ms; τslow = 929.9 ± 325.1 ms). Accordingly, a stimulation protocol to measure slow inactivation of BTX-yne-modified channels requires conditioning pulses >> 1,000 ms in duration to ensure that equilibrium is achieved between successive current recordings.

The effect of each toxin on slow inactivation was examined using the stimulation protocol shown in Figure 3A (inset). A conditioning pulse was applied from −140 to 0 mV in 5 mV increments over a 6,500-ms interval (Δt = 6.5 × τslow). Following the pre-pulse, a 150-ms test pulse to 0 mV measured channel availability. In unmodified channels, a brief hyperpolarizing step (10 ms) to −120 mV was included to recover from fast inactivation (not shown in Figure 3A, see Figure S7). The addition of this step pulse is both unnecessary and problematic in recordings of toxin-modified channels—the former because all four compounds block fast inactivation, the latter stemming from the large and variable tail currents, which interfere with measurements in the test pulse.

**Table 1. Boltzmann fit and affinity parameters for untreated and toxin-modified rNaV1.4**

<table>
<thead>
<tr>
<th>Toxin</th>
<th>V1/2 act (mV)</th>
<th>ΔV1/2 act (mV)</th>
<th>k (μM)</th>
<th>EC50 (nM)</th>
<th>EC50 (from SSI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>−28.9 ± 0.3</td>
<td>−</td>
<td>6.5 ± 0.3</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>BTX-B</td>
<td>−74.3 ± 0.6</td>
<td>−45.4 ± 0.7</td>
<td>4.8 ± 0.5</td>
<td>2074 ± 768</td>
<td>2237 ± 1120</td>
</tr>
<tr>
<td>BTX-yne</td>
<td>−67.5 ± 0.2</td>
<td>−38.6 ± 0.4</td>
<td>4.5 ± 0.2</td>
<td>756 ± 43</td>
<td>795 ± 113</td>
</tr>
<tr>
<td>BTX-B-Hx</td>
<td>−78.4 ± 0.1</td>
<td>−49.5 ± 0.3</td>
<td>2.0 ± 0.1</td>
<td>491 ± 26</td>
<td>494 ± 37</td>
</tr>
<tr>
<td>BTX-yne</td>
<td>−71.4 ± 0.3</td>
<td>−42.7 ± 0.4</td>
<td>6.5 ± 0.3</td>
<td>130 ± 21</td>
<td>189 ± 59</td>
</tr>
</tbody>
</table>

Cells were stepped from a holding potential of −120 to −70 mV for varying lengths of time, Δt. Following this conditioning pulse, the extent to which channels inactivate as a function of Δt was measured in a test pulse to 0 mV (Figure S6) (Cummins et al., 1998). BTX-B was employed as a negative control for these experiments, as channels bound to this compound remain conducting irrespective of the time period of the conditioning or test pulses. Using this stimulation protocol, cells treated with 10 μM BTX-yne display a clear exponential decay of peak current as a function of the conditioning pulse length. Fitting these data for BTX-yne to two exponentials gives a time constant, τ, for each process (τfast = 90.8 ± 3.8 ms; τslow = 929.9 ± 325.1 ms). Accordingly, a stimulation protocol to measure slow inactivation of BTX-yne-modified channels requires conditioning pulses >> 1,000 ms in duration to ensure that equilibrium is achieved between successive current recordings.

The effect of each toxin on slow inactivation was examined using the stimulation protocol shown in Figure 3A (inset). A conditioning pulse was applied from −140 to 0 mV in 5 mV increments over a 6,500-ms interval (Δt = 6.5 × τslow). Following the pre-pulse, a 150-ms test pulse to 0 mV measured channel availability. In unmodified channels, a brief hyperpolarizing step (10 ms) to −120 mV was included to recover from fast inactivation (not shown in Figure 3A, see Figure S7). The addition of this step pulse is both unnecessary and problematic in recordings of toxin-modified channels—the former because all four compounds block fast inactivation, the latter stemming from the large and variable tail currents, which interfere with measurements in the test pulse.

**Table 1. Boltzmann fit and affinity parameters for untreated and toxin-modified rNaV1.4**

<table>
<thead>
<tr>
<th>Toxin</th>
<th>V1/2 act (mV)</th>
<th>ΔV1/2 act (mV)</th>
<th>k (μM)</th>
<th>EC50 (nM)</th>
<th>EC50 (from SSI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>−28.9 ± 0.3</td>
<td>−</td>
<td>6.5 ± 0.3</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>BTX-B</td>
<td>−74.3 ± 0.6</td>
<td>−45.4 ± 0.7</td>
<td>4.8 ± 0.5</td>
<td>2074 ± 768</td>
<td>2237 ± 1120</td>
</tr>
<tr>
<td>BTX-yne</td>
<td>−67.5 ± 0.2</td>
<td>−38.6 ± 0.4</td>
<td>4.5 ± 0.2</td>
<td>756 ± 43</td>
<td>795 ± 113</td>
</tr>
<tr>
<td>BTX-B-Hx</td>
<td>−78.4 ± 0.1</td>
<td>−49.5 ± 0.3</td>
<td>2.0 ± 0.1</td>
<td>491 ± 26</td>
<td>494 ± 37</td>
</tr>
<tr>
<td>BTX-yne</td>
<td>−71.4 ± 0.3</td>
<td>−42.7 ± 0.4</td>
<td>6.5 ± 0.3</td>
<td>130 ± 21</td>
<td>189 ± 59</td>
</tr>
</tbody>
</table>
Using the protocol shown in Figure 3A (inset), absent a hyperpolarizing step to recover from fast inactivation, the inactivation response of unmodified channels (black circles) displays sigmoidal behavior (Figure 3A). When a 10-ms hyperpolarizing step to −120 mV is included to enable channels to recover from fast inactivation, the voltage dependence of SSI is shallower (i.e., larger k value) and shifted considerably to higher potential, and normalized current (I/I_0) never reaches zero (open circles). Channels modified by 10 μM BTX-B do not inactivate in either the conditioning or the test pulse (Figures 3A and 3B, red triangles), analogous to BTX (Wang and Wang, 1994, 1996). By contrast, channels modified with 10 μM BTX-yne exhibit a clear voltage-dependent slow inactivation with current decaying to a steady-state value that is ~60% of the peak current, I_0 (Figures 3A and 3B, blue squares). Two principal conclusions follow from these data: 1) ~80% of unmodified channels enter a slow-inactivated state (open circles, Figure 3C); 2) BTX-yne-modified channels are able to undergo slow inactivation. The latter finding is in marked contrast to channels treated with BTX-B, for which both fast and slow inactivation are entirely inhibited.

To obtain additional support for the above conclusions, we used an amended stimulation protocol to measure fast inactivation (Figure 3C, inset). In these experiments, a short conditioning pulse (10 ms) only allows time for channels to undergo fast inactivation. Accordingly, application of 10 μM BTX-yne to cells under this protocol gives no evidence of inactivation (Figure 3C). In fact, these data look indistinguishable from the inactivation plots of other BTX derivatives (Figure 2B). These results confirm that BTX-yne prevents fast inactivation and that BTX-yne-modified channels are able to access the slow-inactivated state.

**BTX derivatives differentially affect bacterial Na_V inactivation**

Our findings with NaV1.4 and NaV1.5 implicate BTX-yne as an inhibitor of fast but not slow inactivation. To interrogate further the activity of this toxin derivative, we have examined the influence of BTX-yne as well as BTX, BTX-B, and BTX-Hx on Na_\text{Vs} that lack a fast inactivation gate. Bacterial Na_\text{s} (BacNa_\text{s}) form as homotetramers of non-covalently associated domains, analogous to voltage-gated K_+ channels, and lack the DIII–DIV linker and IFM particle (or a functional equivalent) responsible for fast inactivation in eukaryotic channels (Payandeh and Minor Jr., 2015; Lenaues et al., 2017; Payandeh, 2018). Thus, only one form of inactivation is operative in BacNa_\text{s}.

We initially examined Na_Bh1 (NachBac), a well-studied BacNa_\text{V} that is sensitive to BTX (Finol-Urdaneta et al., 2019). In line with our work and previous reports of BTX activity against eukaryotic Na_\text{s}, toxin application (10 μM) stabilized the Na_Bh1 open state and shifted the activation voltage dependence toward hyperpolarizing potentials (Figures S8A, S8B, and S8D). Unlike eukaryotic Na_\text{s}, however, Na_Bh1 proved surprisingly insensitive to BTX-B (Figures S8C and S8D). An alternative BacNa_\text{V}, Na_Ae1, from Alkalimnicola ehrlichii (Arrigoni et al., 2016) bearing a mutation in the cytosolic C-terminal domain “neck region” (R264E) to enable functional studies (Shaya et al., 2014) was only modestly influenced by BTX (Figures S8E–S8G). To understand these unexpected differences in toxin activity, we examined sequence alignments of the inner pore S6 helices of select bacterial and eukaryotic channels (Figure S8H). This analysis identified a serine residue in the Na_Ae1 inner pore that appears as a glycine in Na_Bh1 and in three of the four S6 helices in eukaryotic Na_\text{s} (Figures S8H–S8I). Mutation of S225 to glycine (S225G) in Na_Ae1 R264E yields a channel that is sensitive to both BTX and BTX-B (Figures 4A–4C and S8J–S8K) as well as BTX-Hx and BTX-yne (Figures 4D and 4E). All four compounds cause substantial hyperpolarizing shifts in activation and inhibit channel inactivation (Figures 4A–4F). Electrophysiology recordings to measure steady-state inactivation confirm that BTX, BTX-B, and BTX-Hx significantly block inactivation of Na_Ae1 S225G-R264E (Figures 4B–4D and 4G). In marked contrast, Na_Ae1 S225G-R264E channels exposed to BTX-yne show a strong SSI response, as noted in unmodified channels (Figure 4G). This result offers compelling evidence that BTX-yne, unlike the other three compounds, is ineffective at blocking inactivation in this mutant BacNa_\text{V}.
DISCUSSION

We have synthesized BTX and three C20-ester derivatives and characterized these compounds by whole-cell electrophysiology against rNaV1.4. All four compounds significantly hyperpolarize V_{1/2} of activation, albeit with some variation in the magnitude of this effect (see Table 1). We have examined different protocols to measure the influence of non-saturating toxin concentrations on channel activity and have established that these methods give similar outputs (Li et al., 2002; Wang and Wang, 2017). Concentration-response measurements quantify the relative activity of BTX, BTX-B, BTX-\textsuperscript{2Hx}, and BTX-yne. The potencies of BTX, BTX-B, and BTX-\textsuperscript{2Hx}, the three derivatives that eliminate fast and slow inactivation, track with cLogP values for these compounds (cLogP: 3.22, 4.12, 4.47; EC\textsubscript{50} for V_{1/2} activation: 2074 nM, 756 nM, 491 nM, respectively).\textsuperscript{23} This trend is consistent with toxin binding at site II, as the inner pore of the channel is a receptor for hydrophobic molecules. To our knowledge, this work represents the first systematic analysis of toxin potency against a recombinant Na\textsubscript{V} subtype.

The influence of BTX on channel inactivation is pronounced, as channels remain persistently conducting, unable to conformationally rearrange in a manner that stops the flow of ions (Kholodov, 1985; Brown, 1988). Our findings demonstrate that BTX-B and BTX-\textsuperscript{2Hx}, like the parent toxin, block fast and slow inactivation mechanisms. Channels modified by BTX-yne are also prevented from rapidly inactivating. BTX-yne binding, however, does not preclude channels from undergoing a slow, voltage-dependent transition to a non-conducting state. Slow inactivation is thought to involve extensive protein conformational changes, but a structural understanding of this process is still lacking (Cummins and Sigworth, 1996; Featherstone et al., 1996; Hayward et al., 1997; Vedantham and Cannon, 1998; Vilin et al., 2001; Hilber et al., 2002; Wang et al., 2003; Webb et al., 2009). Toxin derivatives such as BTX-yne should aid studies of this intriguing gating mechanism.

Additional evidence that BTX-yne is unable to block slow inactivation follows from investigations with the bacterial sodium channel mutant, Na\textsubscript{V}Ae1-S225G-R264E. As with other BacNa\textsubscript{s}s, Na\textsubscript{V}Ae1 S225G-R264E lacks a fast inactivation particle and, thus, has only one mechanism for transitioning from an open to a non-conducting state. Electrophysiological recordings of Na\textsubscript{V}Ae1 S225G-R264E following treatment with BTX, BTX-B, or BTX-\textsuperscript{2Hx} display essentially equivalent effects in which all three compounds induce a large hyperpolarizing shift in activation voltage dependence and block inactivation (Figures 4B–4D, 4F, and 4G). These
data are similar to results with eukaryotic channels, NaV1.4 and 1.5. By contrast, NaV_{AE1} S225G-R264E channels modified by BTX-yne, for which activation voltage is significantly hyperpolarized, can enter a non-conducting state (Figures 4E–4G). The inability of BTX-yne to block inactivation in NaV_{AE1} S225G-R264E supports our conclusion that this same compound is incapable of blocking slow inactivation in mammalian NaVs.

A simplified four-state model is sufficient to appreciate the differences between BTX and BTX-yne on NaV gating (for a recent, comprehensive discussion on NaV gating, see Catterall et al., 2020). Closed channels, unmodified by toxin, can enter either an open (conductive) or slow-inactivated (non-conductive) state (Figure 5A). The degree to which either pathway is favored is voltage-dependent. Open channels quickly become non-conducting through a mechanism of fast inactivation and a secondary inactivation pathway that is kinetically slower. BTX, as well as BTX-B and BTX-CDE-Hx, facilitates the closed→open transition and blocks both inactivation processes; thus, channels open upon membrane depolarization at more negative potentials and remain conductive.

As with BTX, BTX-yne influences the steady-state population of closed*→open* channels and inhibits fast inactivation (note: * reflects BTX-yne-bound channels, Figure 5B). Channel inactivation, however, through closed*→slow-inactivated* (i.e., closed-state inactivation) and open*→slow-inactivated* transitions, is not blocked by BTX-yne. The combination of 1) inhibition of fast inactivation and 2) slow conversion from the open*→slow-inactivated* state relative to the time duration of the conditioning pulse (150 ms) in the SSI protocol gives rise to the unusual U-shaped SSI curve for BTX-yne-modified NaVs (see Figure 2B). As conditioning pulses in the SSI protocol are stepped to higher depolarizing potentials, the fraction of channels transitioning from closed*→slow inactivated* first increases in a voltage-dependent manner, leading to the downstroke of the current versus voltage response, which reaches a nadir at −70 mV. Conditioning depolarizations > −70 mV result in fewer channels undergoing closed-state inactivation, leaving a larger number of channels available to activate. Open* channels cannot fast inactivate and do not have sufficient time to slow inactivate* given the 150-ms duration of the conditioning pulse (recall that τ_{slow} = 930 ms, Figure S6). As such, an increase in current is measured at higher step potentials (the upstroke of the U-shaped curve). This interpretation is consistent with the absence of a U-shaped SSI curve for BTX-yne-modified NaVs with longer (6,500 ms) conditioning pulses (Figure 3A).

For a derivative of BTX to function equivalently to the natural product, the full pentacyclic steroidal core as well as an ester substituent at C20 are necessary. Studies with semi-synthetic BTX derivatives demonstrate that removal of the C20-ester results in compounds that lose considerable potency (Figure S9). Reduction in the B-ring double bond in BTX affords a toxin derivative that hyperpolarizes V_{1/2}, but no longer completely blocks inactivation. Synthetic analogues of the AB-ring of BTX containing pendant amine groups antagonize BTX binding, but do not hyperpolarize V_{1/2} or eliminate inactivation (Schow et al., 1997). Additionally, analogues of the CDE-ring of BTX act as reversible NaV inhibitors despite binding in the inner pore of the channel (Toma et al., 2016). Like these compounds, the enantiomer of BTX also functions as a channel antagonist with a binding site that overlaps that of the natural product (Logan, 2016).

Collectively, our findings reveal that the intrinsic rigidity of BTX, including the heteroaryl C20-ester substituent, is critical to agonist function. Modified forms of the toxin that have additional degrees of flexibility (e.g., reduced B-ring double bond, CDE-ring analogues, C20-hexynoate ester) may adopt binding poses that either block ion conduction or enable the channel to enter a partially inactivated conformational state. BTX-yne is the first small molecule tool compound that selectively eliminates NaV fast, but not slow, inactivation. Such a reagent should enable research efforts to explore channel dynamics and the mechanism of this latter process.
CONCLUSION

Our studies to examine structure-activity relationships of BTX and a small collection of ester-modified toxins reveal the complex allosteric properties of these ligands on NaVs function. This work provides a complete electrophysiological characterization of toxin affinity and the effects of toxin binding on both channel activation and inactivation. We have identified a BTX ester derivative, BTX-yne, that affects fast and slow inactivation processes in a manner that is different from BTX and from any other small molecule or protein toxin modulator of NaVs. Future studies will aim to shed light on the molecular details of BTX and BTX-yne binding to the channel. Such insights are expected to inform the rational design of small molecule NaV modulators that offer precise control of gating.

SIGNIFICANCE

NaVs are obligatory protein complexes for generating electrical signals in neuronal cells. These molecular machines operate through complex mechanisms in order to regulate the flow of sodium ions across cellular membranes. Understanding how NaVs function and how small molecules that target these channels modulate ionic conduction will inform efforts in pharmaceutical research to design precision therapeutics for treating NaV-related pathologies.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
  - Lead contact
  - Materials availability
  - Data and code availability
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
  - CHO and HEK cell cultures
- METHOD DETAILS
  - Transient transfection of NaV plasmids
  - Toxin quantification and storage
  - Electrophysiology
  - Chemistry: general experimental
  - Synthetic experimental procedures
- QUANTIFICATION AND STATISTICAL ANALYSIS

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.chembiol.2021.12.003.

ACKNOWLEDGMENTS

Partial support of this work has come from the National Institutes of Health (R01 NS045684) and from a generous gift from Amgen, Inc. T.M.G.M. was supported by a fellowship from the Center for Molecular Analysis and Design (CMAD) at Stanford University. C.E.G. is grateful for support from the Stanford ChEM-H Chemistry/Biology Interface Predoctoral Training Grant Program and the NIGMS of the NIH under Award Number T32GM120007 and to Abbott Laboratories for a Stanford Graduate Fellowship. This work was supported by grants NIH-NHLBI R01-HL080050 and NIH-NIDCD R01-DC007664 to D.L.M. and an American Heart Association postdoctoral fellowship to F.A.-A.

AUTHOR CONTRIBUTIONS

T.M.G.M. and J.D. originally conceived of the project. T.M.G.M., F.A.-A., and C.E.G. performed research; all authors were involved in analyzing the data. T.M.G.M. and J.D. wrote the paper with input from C.E.G., F.A.-A., and D.L.M.

DECLARATION OF INTERESTS

J.D. is a cofounder, executive board member, and holds equity shares in Si-teOne Therapeutics, Inc., a start-up company interested in developing subtype-selective modulators of NaVs.

Received: February 22, 2021
Revised: September 14, 2021
Accepted: November 29, 2021
Published: December 27, 2021

REFERENCES


Cell Chemical Biology Article


### KEY RESOURCES TABLE

<table>
<thead>
<tr>
<th>REAGENT or RESOURCE</th>
<th>SOURCE</th>
<th>IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Chemicals, peptides, and recombinant proteins</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dulbecco’s Modified Eagle Medium (DMEM), high glucose</td>
<td>GIBCO, Grand Island, NY</td>
<td>Cat# 11965-118</td>
</tr>
<tr>
<td>Cosmic calf serum</td>
<td>HyClone, Logan, UT</td>
<td>Cat# SH3008703</td>
</tr>
<tr>
<td>Penicillin-streptomycin (10,000 U/mL)</td>
<td>GIBCO, Grand Island, NY</td>
<td>Cat# 15140122</td>
</tr>
<tr>
<td>Trypsin-EDTA 1X Solution</td>
<td>Invitrogen, Carlsbad, CA</td>
<td>Cat# 59417C</td>
</tr>
<tr>
<td>Batrachotoxin (BTX)</td>
<td>This study</td>
<td>N/A</td>
</tr>
<tr>
<td>Batrachotoxin-B (BTX-B)</td>
<td>This study</td>
<td>N/A</td>
</tr>
<tr>
<td>Batrachotoxin-Hx (BTX-Hx)</td>
<td>This study</td>
<td>N/A</td>
</tr>
<tr>
<td>Batrachotoxin-yne (BTX-yne)</td>
<td>This study</td>
<td>N/A</td>
</tr>
</tbody>
</table>

#### Experimental Models: Cell Lines

<table>
<thead>
<tr>
<th>RESOURCE or RESOURCE SOURCE IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chinese hamster ovary (CHO) cells</td>
</tr>
<tr>
<td>Human embryonic kidney (HEK293) cells</td>
</tr>
</tbody>
</table>

#### Recombinant DNA

<table>
<thead>
<tr>
<th>RESOURCE or RESOURCE SOURCE IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td>rSkM1 clone  ( \mu_1-2 + pZem228-rNa_y1.4 )</td>
</tr>
<tr>
<td>pcDNA3.1(+)HNa_y1.5</td>
</tr>
<tr>
<td>pIRES-EGFP-NaVAe1</td>
</tr>
<tr>
<td>pTRACER-CMV2-NaVBh1</td>
</tr>
</tbody>
</table>

#### Software and Algorithms

<table>
<thead>
<tr>
<th>RESOURCE or RESOURCE SOURCE IDENTIFIER</th>
</tr>
</thead>
</table>

#### Other

<table>
<thead>
<tr>
<th>RESOURCE or RESOURCE SOURCE IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Axopatch-200A Amplifier</td>
</tr>
<tr>
<td>Digidata 1322B Digitizer</td>
</tr>
<tr>
<td>Borosilicate glass micropipettes</td>
</tr>
</tbody>
</table>

### RESOURCE AVAILABILITY

#### Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Justin Du Bois (jdubois@stanford.edu).  

#### Materials availability

Toxin derivatives generated in this study are limited in supply due to their difficulty of synthesis and toxicity, and require a material transfer agreement for distribution. Plasmids and cell lines can be requested by contacting the lead contact.
Data and code availability
- All data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

CHO and HEK cell cultures
Chinese hamster ovary (CHO) cells were obtained from Prof. Jon Sack (UC Davis) and were not further authenticated. Chinese hamster ovary and human embryonic kidney (HEK) cell cultures were prepared as described previously (Andresen and Du Bois, 2009; Shaya et al., 2014). Briefly, CHO cells were grown in DMEM (GIBCO, Grand Island, NY) supplemented with 10% cosmic calf serum (HyClone, Logan, UT) and 100 IU/mL penicillin/streptomycin (GIBCO). Cells were kept in a 5% carbon dioxide, 96% relative humidity incubator at 37 °C and passed every ~3 days. Passaging of cells was accomplished by aspiration of media, washing with phosphate-buffered saline, treatment with 1 mL trypsin-EDTA (0.05%, Invitrogen, Carlsbad, CA) for ~5 min until full dissociation of cells from the plate surface was observed and dilution with 4 mL of growth medium. Approximately 250 μL of this suspension was then diluted in 10 mL of growth medium in a new 10 cm plate. Human embryonic kidney cells (HEK293) were purchased from ATCC (CRL-1573) and were grown at 37 °C under 5% CO₂, in a Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 10% l-glutamine, and antibiotics (100 IU/mL penicillin and 100 mg/mL streptomycin) (University of California, San Francisco Cell Culture Facility). The sex of cell line is not determined.

METHOD DETAILS

Transient transfection of Na₉ plasmids
Transfection with a pZem228 vector containing the full-length cDNA coding for the α-subunit of rNaᵥ1.4 was accomplished using the calcium phosphate precipitation method. Transfection with a pcDNA3.1(+) vector containing the full-length cDNA coding for the α-subunit of hNaᵥ1.5 was accomplished using Lipofectamine LTX. Transfection with a pIRE2-EGFP vector containing the full-length cDNA coding for NaᵥAE1 or NaᵥBh1 constructs was accomplished using Lipofectamine 2000. Cotransfection with EGFP was employed so that fluorescence of EGFP could be used as a marker of transfection efficiency.

Toxin quantification and storage
BTX derivatives were quantified by ¹H NMR spectroscopy on a 600 MHz Varian Inova spectrometer. 1,3-benzodioxole was employed as an internal standard. Spectra were acquired with a relaxation delay time (d1) of 20 s and an acquisition time (at) of 10 s. Quantitation was determined by comparison of the integrations for the toxin and the internal standard of a known concentration. Toxin stock solutions prepared in dimethyl sulfoxide (4 mM) were kept at −20 °C and diluted with external solution before recording.

Electrophysiology
Sodium currents were measured using the patch-clamp technique in the whole-cell configuration with an Axopatch-200b amplifier (Axon Instruments, Union City, CA), as previously described by Moran and coworkers (Moran et al., 2003). Borosilicate glass micropipettes (Sutter Instruments, Novato, CA) were fire-polished to a tip diameter yielding a resistance of 1.2–4.5 MΩ in the working solutions. The micropipette was filled with 40 mM NaF, 1 mM EDTA, 20 mM HEPES, 125 mM CsCl. The external solution had the following composition: 180 mM NaCl, 2 mM CaCl₂, 20 mM HEPES. The pH of the solutions was adjusted to pH 7.4 with 50 wt% aqueous CsOH.

The output of the patch-clamp amplifier was filtered with a built-in low-pass, four-pole Bessel filter having a cutoff frequency of 10 kHz and sampled at 100 kHz. The membrane was kept at a holding potential of −100 mV. Pulse stimulation and data acquisition used 16-bit D-A and A-D converters (Axon Instruments DigiData 1322A) controlled with the pClamp software (Axon Instruments). Leak currents were subtracted using a standard P/4 protocol of the same polarity. Access resistance was always <4 MΩ and the cell capacitance was between 4 and 20 pF, as measured by the compensating circuit of the feedback amplifier. Peak currents were generally between 1 and 5 nA. The series resistance was typically compensated 80%. All measurements were done at ambient temperature (20–22 °C). Recordings were made at least 5 min after establishing the whole-cell and voltage-clamp configuration to allow for stabilization of the voltage-dependent properties of the channels.

Cells were modified by toxin after stabilization of whole-cell voltage parameters and control measurements. 0.5 mL of toxin solution was applied over 60 s. Toxin binding was promoted via 2,000 pulses to 0 mV from a hold potential of −100 mV (2 Hz cycling) before measurements were performed.

Chemistry: general experimental
Unless stated otherwise, all reactions were performed at ambient temperature in flame-dried or oven-dried glassware under an argon or nitrogen atmosphere using dry, deoxygenated solvents (distilled or passed over a column of activated alumina). Air- and moisture-sensitive liquids and solutions were transferred via syringe or stainless steel cannula. Organic solutions were concentrated under reduced pressure (~15 Torr) by rotary evaporation. Commercially available reagents were used as received. Triethylamine was
distilled from calcium hydride immediately prior to use. Reactions requiring external heat were modulated to specified temperatures using an IKAmag temperature controller. Thin-layer chromatography (TLC) was performed using E. Merck silica gel 60 F254 pre-coated plates (250 nm) and visualized by UV fluorescence quenching, potassium permanganate, aqueous ceric ammonium molybdate, or p-anisaldehyde staining. Silicycle SiliaFlash P60 Academic Silica gel (particle size 40–63 nm) was used for flash chromatography. In select cases as indicated, NH4OH pre-treated silica gel was used for chromatographic separation as described previously (Logan, 2016). Briefly, preparation of NH4OH pre-treated silica gel was accomplished by slurring silica gel with concentrated aqueous NH4OH in a large crystallization dish, followed by evaporation of water by heating at 50°C on a hot plate overnight.

1H and 13C NMR spectra were recorded on a Varian Inova 400 (400 MHz and 100 MHz, respectively), a Varian Inova 500 (500 MHz and 125 MHz, respectively), or a Varian Inova 600 (600 MHz and 150 MHz, respectively) and are reported in terms of chemical shift relative to residual CHCl3 (in CDCl3, δ 7.26 and δ 77.16, respectively) or C6H6 (in C6D6, δ 7.16 and δ 128.39, respectively). Data for 1H NMR spectra are reported as follows: chemical shift (δ, ppm), multiplicity (s, singlet; d: doublet; t, triplet; q, quartet; m, multiplet), coupling constant (Hz), integration. Data for 13C are reported in terms of chemical shift (δ, ppm). Infrared spectra were recorded as thin films using NaCl salt plates on a Thermo-Nicolet 300 FT-IR and are reported in frequency of absorption. High-resolution mass spectra were obtained from the Vincent Coates Foundation Mass Spectrometry Laboratory at Stanford University. Samples were analyzed by liquid chromatography-electrospray ionization mass spectrometry on a Waters Acquity UPLC and Thermo Fisher Exactive Mass Spectrometer scanning m/z 100-1,000. The LC mobile phase was 100% methanol and the flow rate was 0.175 mL/min.

**Caution**

Batrachotoxin (BTX) and synthetic derivatives thereof are (or may be) extremely potent neurotoxins (LD50 = 1–2 μg/kg; mouse, subcutaneous) (Tokuyama et al., 1969) and should be handled with great care. The compounds are skin permeable and may cause paralysis or death if contact with skin occurs. Proper personal protective equipment should be worn at all times.

**Synthetic experimental procedures**

**Cyclohexanecarboxylic (ethyl carbonic) anhydride**

To an ice-cold solution of cyclohexane-carboxylic acid (548 mg, 4.27 mmol) in 21 mL of benzene was added sequentially Et3N (410 μL, 4.27 mmol) and ethyl chloroformate (600 μL, 4.27 mmol). The solution was stirred at 0°C for 30 min. Following this time, the reaction mixture was transferred to a separatory funnel with 30 mL of Et2O and 30 mL of 1.0 M aqueous NaOH. The organic layer was collected and the aqueous layer was extracted with 3 x 50 mL of Et2O. The combined organic extracts were dried over MgSO4, filtered, and concentrated under reduced pressure to a yellow oil. Purification of this material by chromatography on silica gel (20% Et2O/hexanes) afforded cyclohexanecarboxylic (ethyl carbonic) anhydride as a clear oil (692 mg, 81%).

**1H NMR (500 MHz, CDCl3)**

δ 4.31 (q, J = 7.1 Hz, 2H), 2.43 (tt, J = 11.3, 3.7 Hz, 1H), 1.98 (d, J = 13.2 Hz, 2H), 1.81–1.75 (m, 2H), 1.68–1.61 (m, 2H), 1.56–1.43 (m, 2H), 1.36 (t, J = 7.2 Hz, 3H), 1.32–1.19 (m, 2H) ppm

**Hept-6-ynecarboxylic (ethyl carbonic) anhydride**

To an ice-cold solution of hept-6-yne carboxylic acid (299 mg, 2.37 mmol) in 12 mL of benzene was added sequentially Et3N (230 μL, 2.37 mmol) and ethyl chloroformate (330 μL, 2.37 mmol). The solution was stirred at 0°C for 30 min. Following this time, the reaction mixture was transferred to a separatory funnel with 30 mL of Et2O and 30 mL of 1.0 M aqueous NaOH. The organic layer was collected and the aqueous layer was extracted with 3 x 50 mL of Et2O. The combined organic extracts were dried over MgSO4, filtered, and concentrated under reduced pressure to a yellow oil. Purification of this material by chromatography on silica gel (10% EtOAc/hexanes) afforded the desired product as a clear oil (368 mg, 78%).

**TLC Rf = 0.55 (25% EtOAc/hexanes)**

**1H NMR (400 MHz, CDCl3)**

δ 4.32 (q, J = 7.1 Hz, 2H), 2.43 (tt, J = 11.3, 3.7 Hz, 1H), 1.98 (d, J = 13.2 Hz, 2H), 1.81–1.75 (m, 2H), 1.68–1.61 (m, 2H), 1.56–1.43 (m, 2H), 1.36 (t, J = 7.2 Hz, 3H), 1.32–1.19 (m, 2H) ppm.
To a solution of batrachotoxinin A (1.6 mg, 3.8 μmol) in 2.0 mL of anhydrous benzene were sequentially added Et₃N (150 μL, 1.1 mmol, 289 equiv) and cyclohexanecarboxylic (ethyl carbonic) anhydride (10 mg, 49.9 μmol, 13.1 equiv). The reaction mixture was stirred at 45 °C for 18 h. Following this time, all volatiles were removed under reduced pressure to give a pale yellow residue. This material was transferred to a 16 × 125 mm test tube with 4 mL of CHCl₃. The solution was cooled in an ice bath and 4.0 mL of ice-cold 0.1 M aqueous HCl was added slowly. With the aid of a glass pipet, the layers were mixed, the CHCl₃ layer was carefully removed, and the aqueous layer was extracted with 1 × 4.0 mL of CHCl₃. The combined organic fractions were discarded. The pH of the aqueous layer was then adjusted to 10 with 2 mL of 1.0 M aqueous NH₄OH, and the solution was extracted with 3 × 4 mL of CHCl₃. The combined organic fractions were dried over Na₂SO₄, filtered, and concentrated under reduced pressure to a colorless residue. Purification of this material by chromatography on NH₄OH pre-treated silica gel (gradient elution: 30% EtOAc/pentane → 70% EtOAc/pentane with 1% v/v Et₃N) afforded BTX-²Hx as a white solid (1.3 mg, 65%).

TLC R<sub>f</sub> = 0.41 (10% MeOH/CH₂Cl₂ on NH₄OH pre-treated silica gel plates)

<sup>1</sup>H NMR (600 MHz, CDCl₃) δ 6.18-6.14 (m, 1H), 5.82 (s, 1H), 5.67 (q, J = 6.5 Hz, 1H), 3.70 (td, J = 10.7, 3.6 Hz, 1H), 3.64 (dt, J = 13.4, 4.5 Hz, 1H), 3.52 (ddd, J = 13.2, 9.0, 4.0 Hz, 1H), 3.17 (d, J = 17.6 Hz, 1H), 2.93–2.86 (m, 1H), 2.79 (d, J = 14.0 Hz, 1H), 2.59 (d, J = 13.2 Hz, 1H), 2.53 (d, J = 14.1, 1H), 2.44 (ddd, J = 18.9, 4.7, 2.2 Hz, 1H), 2.34 (s, 3H), 2.31–2.19 (m, 2H), 2.16 (dd, J = 13.0, 10.6 Hz, 3H), 1.92 (dd, J = 13.8, 11.2 Hz, 4H), 1.82 (td, J = 12.6, 3.9 Hz, 1H), 1.77 (s, 1H), 1.75 (s, 3H), 1.64 (q, J = 17.0, 14.5 Hz, 4H), 1.54 (s, 3H), 1.48 (dd, J = 9.3, 4.0 Hz, 1H), 1.41 (d, J = 6.4 Hz, 3H), 1.38–1.30 (m, 1H), 1.07 (s, 1H), 0.90 (s, 3H) ppm

HRMS (ESI<sup>+</sup>) calcd for C₃₁H₄₅NO₆ 527.3247 found 528.3313 (M+H<sup>+</sup>)

To a solution of batrachotoxininin A (1.3 mg, 3.1 μmol) in 2.0 mL of anhydrous benzene were sequentially added Et₃N (100 μL, 730 μmol, 235 equiv) and hept-6-ynecarboxylic (ethyl carbonic) anhydride (10 mg, 50.4 μmol, 16.3 equiv). The reaction mixture was stirred at 45 °C for 18 h. Following this time, all volatiles were removed under reduced pressure to give a pale yellow residue. This material was transferred to a 16 × 125 mm test tube with 4 mL of CHCl₃. The solution was cooled in an ice bath and 4.0 mL of ice-cold 0.1 M aqueous HCl was added slowly. With the aid of a glass pipet, the layers were mixed, the CHCl₃ layer was carefully removed, and the aqueous layer was extracted with 1 × 4.0 mL of CHCl₃. The combined organic fractions were discarded. The pH of the aqueous layer was then adjusted to 10 with 2 mL of 1.0 M aqueous NH₄OH, and the solution was extracted with 3 × 4 mL of CHCl₃. The combined organic fractions were dried over Na₂SO₄, filtered, and concentrated under reduced pressure to a colorless residue. Purification of this material by chromatography on NH₄OH pre-treated silica gel (gradient elution: 30% EtOAc/pentane → 70% EtOAc/pentane with 1% v/v Et₃N) afforded BTX-yne as a white solid (0.9 mg, 56%).

TLC R<sub>f</sub> = 0.39 (25% pentane/EtOAc with 10% v/v Et₃N)
QUANTIFICATION AND STATISTICAL ANALYSIS

Data were analyzed using the Igor Pro 6.37 environment (Wavemetrics). Fit of data is presented as mean ± SD. All experiments are presented as mean ± SEM for \( n \geq 3 \) independent measurements. All statistical details can be found in figure legends and/or table captions.