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1 **Production and Stability of Oxygen-18 Labeled Caribbean Ciguatoxins and Gambierones**

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21

1 **Abstract**

2 Ciguatoxins (CTXs) and gambierones are ladder-shaped polyethers associated with
3 ciguatera poisoning and *Gambierdiscus* spp. Several of these compounds contain carbonyl or
4 hemiketal groups, which have the potential to exchange with ^{18}O -labeled water under acidic
5 conditions. The effects of solvent composition and acid on the rate of exchange and on the
6 stability of the labels at various pH values were assessed to optimize the incorporation of ^{18}O
7 into Caribbean ciguatoxin-1 and -2 (C-CTX1/2), gambierone, and 44-methylgambierone.
8 LC–HRMS results showed that ^{18}O -labeling occurred at the hydroxy group of the hemiketal at
9 C-56 in C-CTX1/2, and at the hydroxy group of the hemiketal at C-4 and the ketone at C-40 in
10 gambierones. Labeling occurred very rapidly (complete in < 30 min) for C-CTX1/2, and more
11 slowly (complete in ca. 16 h) for both gambierones. Labeled C-CTX1/2 was reduced with
12 sodium borohydride to produce ^{18}O -labeled C-CTX3/4. The incorporated ^{18}O labels in the
13 gambierones and C-CTXs were retained in aqueous solvent mixtures under neutral conditions in
14 a short-term stability study, demonstrating that these ^{18}O -labeled toxins have the potential to be
15 used in isotope dilution and metabolism studies.

16 **Keywords:** ciguatoxin, gambierone, LC–HRMS, stable isotope labeling, H_2^{18}O , oxygen-18

17

1 **1. Introduction**

2 Ciguatera poisoning is caused by the consumption of ciguatoxin-contaminated seafood
3 including commercially relevant fish harvested from tropical and sub-tropical regions.
4 Ciguatoxins (CTXs) production has been linked to the benthic dinoflagellate genera
5 *Gambierdiscus* and *Fukuyoa* with toxins moving into marine food webs via herbivory fish.
6 Several precursor Pacific CTXs have been identified in these genera (Chinain et al., 2010), and
7 have been shown to undergo biotransformation into the more toxic CTXs found in fish flesh
8 (Ikehara et al., 2017).

9 Ciguatoxins are large ladder-shaped polyether compounds with molecular masses of
10 1000–1150 Da. They are odourless, tasteless and heat-stable compounds that are potent voltage-
11 gated sodium channel activators, and very low levels are required to induce a toxic effect
12 (Friedman et al., 2008). While no regulatory limit for CTXs has been established, the guidance
13 levels of 0.01 µg/kg CTX1B and 0.1 µg/kg Caribbean CTXs (C-CTXs) in fish flesh suggested by
14 the US Food and Drug Administration has also been endorsed by the European Food Safety
15 Authority (EFSA Panel on Contaminants in the Food Chain, 2010; Food and Agriculture
16 Organization of the United Nations and World Health Organization, 2020; Food and Drug
17 Administration, 2021). Detection of CTXs at these levels using mass spectrometry can be
18 difficult due to significant matrix effects, low recovery, poor ionization efficiency, and in-source
19 fragmentation, which all contribute to poor sensitivity (Harwood et al., 2017; Suzuki et al.,
20 2017).

21 *Gambierdiscus* and *Fukuyoa* spp. produce a wide array of ladder-shaped polyethers
22 including gambierol (Satake et al., 1993a), gambieric acids (Morohashi et al., 2000), maitotoxins
23 (Murata et al., 1994), CTXs (Satake et al., 1996; Satake et al., 1993b) and gambierones

1 (Rodríguez et al., 2015). Gambierone and 44-methylgambierone are members of a class of
2 sulfated polyethers identified across several species of these genera and in *Coolia tropicalis*
3 (Murray et al., 2021; Murray et al., 2020; Tibiriçá et al., 2020). Gambierones exhibit CTX-like
4 effects on sodium channels, although at a much lower potency than CTXs, and have minimal
5 toxicity following intraperitoneal injection in mouse bioassays (Boente-Juncal et al., 2019;
6 Murray et al., 2021; Murray et al., 2020; Rodríguez et al., 2015). While the diversity and
7 distribution of gambierones across *Gambierdiscus* and other benthic dinoflagellate genera is not
8 fully understood, the presence of known gambierones in several species suggests they have
9 potential to be used as biomarkers for environmental monitoring of *Gambierdiscus* dominance in
10 coral reef systems (Murray et al., 2020; Yon et al., 2021).

11 LC–MS analysis of complex samples, including fish tissues, can result in performance
12 issues associated with matrix effects and sample preparation that can be improved through the
13 use of isotopically labeled analytes as internal standards (Haddad et al., 2019; Stokvis et al.,
14 2005). Furthermore, isotopically labeled analytes can be used to investigate in vitro and in vivo
15 metabolism, where the use of mass spectrometry can provide accurate metabolite tracing and
16 potentially identify pathways associated with metabolism (Mutlib, 2008). There is limited
17 availability of CTX standards, especially those associated with the Caribbean region. Several
18 algal CTXs can be isolated from toxin-producing *Gambierdiscus* isolates, but this will not be
19 feasible for C-CTXs until C-CTX-producing algae can be identified and brought into culture.
20 With their complex structures and laborious synthesis schemes (Inoue et al., 2006; Sasaki et al.,
21 2021), availability of standards relies on isolation from incurred fish tissue. This limits the
22 possibilities for incorporating isotopic labeling during laboratory synthesis, or by modification of
23 culture conditions such as that used in the production of ¹⁸O-labeled yessotoxins (Yamazaki et

1 al., 2012). At this time, there are no commercially available isotopically labeled CTX standards.
2 An alternative strategy for stable isotope labeling would be to utilize acid-catalyzed oxygen
3 exchange. In this case, carbonyl groups exchange oxygen with water by the reversible formation
4 of hydrates (Theodorou et al., 2014). For example, ¹⁸O-labeling of peptides on their carboxyl-
5 containing amino acid residues has proven to be a successful method for isotopic labeling (Niles
6 et al., 2009).

7 In this study, we investigated the feasibility of oxygen exchange with C-CTXs and
8 gambierones using ¹⁸O-labeled water. The aim of this work was to identify suitable reaction
9 conditions for label incorporation and the extent of labeling for these marine toxins. The labeling
10 kinetics and label stability were analyzed to evaluate their potential for use in analytical and
11 biological studies.

12

13 **2. Material and Methods**

14 2.1. Chemical and Reagents

15 Acetonitrile and formic acid (~98 %) were LC–MS grade from Fisher Scientific (Ottawa,
16 ON, Canada). Ammonium acetate (LC–MS grade, ~98%), sodium borohydride (~98%) and m-
17 aminophenylboronic acid–agarose (mAPBAG) aqueous gel suspension were from Millipore–
18 Sigma (Oakville, ON, Canada). Glass-distilled dichloromethane was from Caledon Laboratories
19 (Georgetown, ON, Canada). Additional MeOH, dichloromethane, and hexane used in
20 preparation of fish materials was HPLC grade from Fisher Scientific (Waltham, MA, USA).
21 H₂¹⁸O (97 atom-% ¹⁸O) was from Cambridge Isotope Laboratories (Tewksbury, MA, USA).
22 Distilled water was ultra-purified to 18.2 MΩ·cm using a Milli-Q water purification system
23 (Millipore–Sigma). All solvent mixtures were prepared by volume. Gambierone (19.9 µg/mL in

1 MeOH) was from CIFGA (Lugo, Spain) and 44-methylgambierone (25 µg/mL in MeOH) was
2 from Cawthron Institute (Nelson, New Zealand).

3 2.2. Preparation of Semi-Purified C-CTX1/2

4 Due to the current lack of reference materials for C-CTXs, semi-purified C-CTX1/2 was
5 prepared from C-CTX-laden fish muscle tissue (*Sphyraena barracuda* and *Scomberomorus*
6 *cavalla*), and a small aliquot of the semi-pure toxin isolate was used in this study. Briefly, fish
7 previously collected near St. Thomas, U.S. Virgin Islands that had been confirmed to be toxic via
8 an in vitro ouabain-veratrine dependent mouse neuroblastoma assay, and the presence of C-
9 CTX-1/2 verified using LC–MS/MS analysis (described elsewhere (Robertson et al., 2014)),
10 were homogenized, combined, and subsequently extracted in MeOH (2 mL/g) three times.
11 Extracts were dried by rotary evaporation at 50 °C, then reconstituted in 80% aqueous MeOH
12 and twice partitioned with hexane (1:1). The methanolic phase was subsequently adjusted to 60%
13 aqueous MeOH and partitioned three times with CH₂Cl₂ and dried at 40 °C by rotary
14 evaporation. The residue was reconstituted in CH₂Cl₂ and loaded onto an open column packed
15 with silica gel (BDH, 120 g, 60 Å, 60–200 µm; VWR, Suwanee, GA, USA) and eluted by
16 gravity. The column was preconditioned with approximately 5 bed volumes of CH₂Cl₂. The C-
17 CTX1/2 pool was eluted in 95:5 CH₂Cl₂–MeOH, dried and then loaded onto a prepacked silica
18 cartridge (5 g/20 mL, Strata® SI-1 Silica, 55 µm, 70 Å; Phenomenex, Torrance, CA, USA) that
19 had been preconditioned with CH₂Cl₂. The C-CTX1/2 was eluted with 4% MeOH in CH₂Cl₂ and
20 dried. The residue was further fractionated by semi-preparative HPLC on a Luna PFP(2) column
21 (150 × 10 mm, 5 µm; Phenomenex, Torrance, CA, USA) at 30 °C with a mobile phase composed
22 of water (A) and methanol (B) with gradient elution (3.0 mL/min) as follows: 0–1 min, 60% B;

1 1–15 min, 60–100% B; 15–22 min, 100% B; followed by re-equilibration at 60% B. Fractions
2 containing C-CTX1/2 were pooled and verified by LC–HRMS analysis (Kryuchkov et al., 2020).

3 2.3. ^{18}O -Exchange Experiments

4 *Experiment 1 (preliminary investigation)*: Aliquots (10 μL) of *S. barracuda* extract, gambierone
5 and 44-methylgambierone were prepared separately by evaporation under N_2 at 35 $^\circ\text{C}$ in glass
6 vials. The residues were each dissolved in 8:4:1 $\text{MeCN-H}_2^{18}\text{O}$ –formic acid (10–50 μL). The
7 solutions were vortex-mixed for 1 min and allowed to stand at ambient temperature for 48 h,
8 then transferred to a vial insert and analyzed by LC–HRMS.

9 *Experiment 2 (labeling kinetics)*: Gambierone and 44-methylgambierone (10 μL of each) were
10 added into the same vial and evaporated under N_2 at 35 $^\circ\text{C}$. An aliquot of *S. barracuda* extract
11 (10 μL) was aliquoted into a separate vial and also evaporated under N_2 at 35 $^\circ\text{C}$. The residues
12 were dissolved in 40:9:1 $\text{MeCN-H}_2^{18}\text{O}$ –formic acid (50–100 μL) and vortex-mixed for 1 min.
13 The solutions were transferred to vial inserts and placed in the LC autosampler at 25 $^\circ\text{C}$ and
14 analyzed repeatedly for 17 h by LC–HRMS.

15 *Experiment 3 (borohydride reduction of ^{18}O -labeled C-CTXs and gambierones)*: Semi-purified
16 C-CTX1/2 (2 μL) and gambierone (10 μL) were evaporated under N_2 at 35 $^\circ\text{C}$ in separate vials
17 and prepared according the procedure described in *Experiment 1* and allowed to stand for 2 h and
18 24 h, respectively. NaBH_4 (1 mg) was added to the solutions and allowed to react for 10 min.
19 The reaction was terminated by addition of 10% formic acid (10 μL) and the solution filtered
20 through a PVDF filter (0.22 μm , Canadian Life Sciences; Peterborough, ON, Canada) at 6010 \times
21 g. The filtrate was transferred to a vial insert and analyzed by LC–HRMS.

22 *Experiment 4 (removal of acid from ^{18}O -labeled C-CTX1/2 by partitioning)*: Semi-purified C-
23 CTX1/2 (2.5 μL) was evaporated under N_2 at 35 $^\circ\text{C}$ and dissolved in 45:45:2 $\text{MeCN-H}_2^{18}\text{O}$ –

1 formic acid (92 μL). The solution was vortex-mixed for 1 min and allowed to stand at ambient
2 temperature for 2 h. The aqueous content of the solution was increased by the addition of H_2^{18}O
3 (20 μL) and transferred to a glass conical test tube. Dichloromethane (CH_2Cl_2) (100 μL) was
4 added and the solution was vortex-mixed for 1 min and centrifuged at $260 \times g$ for 3 min to
5 separate the two layers. The CH_2Cl_2 layer was removed and evaporated under N_2 at 40 $^\circ\text{C}$,
6 dissolved in 100 μL of MeCN, and transferred to a vial insert for LC–HRMS analysis.

7 2.4. Partial Factorial Study of [^{18}O]C-CTX1/2 Labeling Parameters

8 A two-level partial factorial design was applied to the labeling of CTXs by modifying the
9 conditions described in *Experiment 1*. The parameters evaluated included: the proportion of ^{18}O -
10 water, the type and concentration of the acid used, and temperature. A low and a high level was
11 used for each parameter as described (Table 1). Aliquots (10 μL) of *S. barracuda* extract were
12 prepared separately by evaporation under N_2 at 35 $^\circ\text{C}$ in glass vials. The residues were each
13 dissolved in MeCN– H_2^{18}O –acid (10 μL) according to Table 1. The solutions were vortex-mixed
14 for 1 min and allowed to stand at ambient temperature for 48 h, then transferred to a vial insert
15 and analyzed by LC–HRMS to determine the extent of labeling for each trial.

16 2.5. LC–HRMS Analysis

17 Analyses were performed according to Kryuchkov et al. (2020) with some modifications.
18 An Agilent 1200 LC was equipped with a binary pump, temperature-controlled autosampler (10
19 $^\circ\text{C}$) and column compartment (40 $^\circ\text{C}$) (Agilent Technologies, Mississauga, ON, Canada) coupled
20 to a Q Exactive HF Orbitrap mass spectrometer (Thermo Fischer Scientific, Waltham, MA,
21 USA) with a heated electrospray ionization probe (HESI-II). Chromatographic separation was on
22 an F5 column (100 mm \times 2.1 mm, 1.7 μm ; Phenomenex, Torrance, CA, USA) using gradient
23 elution. The mobile phase was composed of water (A) and 95:5 acetonitrile–water (B), each

1 containing 5 mM ammonium acetate (pH 6.8). The gradient employed varied for gambierones
2 and C-CTXs. The gradient for C-CTXs was as follows: 0–18 min, 30–60% B; 18–18.1 min, 60–
3 99% B; 18.1–22 min, 99% B; followed by a 4-min re-equilibration at 30% B. The gradient for
4 gambierones was as follows: 0–18 min, 10–80% B; 18–18.1 min, 80–99% B; 18.1–22 min, 99%
5 B; followed by an 8 min re-equilibration with 10% B. The flow rate was 0.3 mL/min with an
6 injection volume of 5.0 μ L.

7 Full-scan acquisition was performed with a range of m/z 1000–1250 for C-CTXs in
8 positive polarity, and m/z 800–1400 for gambierones with positive and negative polarity
9 switching. The spray voltage of the source was ± 4500 V, with a capillary temperature of 400 $^{\circ}$ C.
10 The sheath and auxiliary gas were set at 45 and 10, respectively, with a max spray current of 100
11 μ A. The probe heater temperature was set at 250 $^{\circ}$ C and the S-Lens RF level was set to
12 maximum (100). The mass resolution was set at 120 000 with an AGC target of 5×10^6 and a
13 maximum injection time of 512 ms per scan.

14 Product-ion spectra were acquired using parallel reaction monitoring (PRM) in positive
15 mode with an isolation window of 1 Da. The mass resolution setting was set at 240 000 with an
16 AGC target of 5×10^6 and a maximum injection time of 512 ms, with a normalized collision
17 energy of 12 for C-CTXs, and a collision energy of 30 eV for gambierones.

18 2.6. Isotope Distribution Analysis

19 Isotopic peak height profiles were collected from LC–HRMS spectra of the protonated or
20 deprotonated molecule ($[M+H]^+$, $[M-H]^-$) as well as sodium ($[M+Na]^+$) or ammonium
21 ($[M+NH_4]^+$) adducts of the compounds of interest. The observed isotopic profiles were used to
22 extract the isotopic composition of oxygen-18 of the CTXs and gambierones. By knowing the
23 identity of the analyzed compounds (their molecular formulae), we established the corresponding

1 isotopic patterns using Fourier-transform-based methods described by Ipsen (2014) and
2 implemented in R package ecipex (<https://CRAN.R-project.org/package=ecipex>). By specifying
3 the number of exchangeable sites for incorporation of oxygen-18, and adopting natural isotopic
4 composition of all other makeup elements in the molecules, the isotopic composition of oxygen
5 at these sites was obtained using partial least-squares fitting. The NRC Isotopic Enrichment
6 Calculator (Mallia et al., 2019) was modified to implement these calculations (Version
7 December 2021; currently available at [https://metrology.shinyapps.io/isotopic-enrichment-](https://metrology.shinyapps.io/isotopic-enrichment-calculator)
8 [calculator](https://metrology.shinyapps.io/isotopic-enrichment-calculator) with source code available from [https://github.com/meijaj/isotopic-enrichment-](https://github.com/meijaj/isotopic-enrichment-calculator)
9 [calculator](https://github.com/meijaj/isotopic-enrichment-calculator)). Parsing mass spectra provides isotopic composition of oxygen at each of the labeling
10 sites and the relative abundances of isotopologues having, in the case of gambierones, 0–2
11 oxygen-18 atoms incorporated. These results were then used to fit the kinetic model of oxygen-
12 18 uptake as described below. The changes in the abundance of label at different locations in the
13 molecule and the total extent of labeling were plotted in SigmaPlot (version 14.0) against time
14 for *Experiment 2*.

15 2.7. Analysis of the Isotopic Label Stability

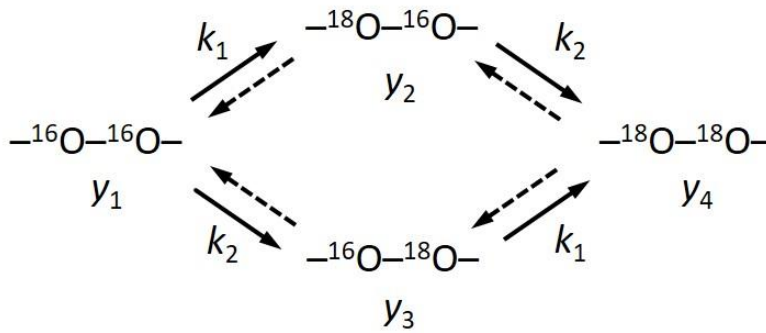
16 C-CTX1/2 (10 μL) was evaporated under N_2 at 35 $^\circ\text{C}$ and reconstituted in 45:45:2
17 MeCN– H_2^{18}O –formic acid (92 μL). The solution was vortex-mixed for 1 min and allowed to
18 stand at ambient temperature for 2 h. The aqueous content of the solution was increased by the
19 addition of H_2^{18}O (20 μL) and transferred to a glass conical test tube. CH_2Cl_2 (100 μL) was
20 added and the solution was vortex-mixed for 1 min and centrifuged at $260 \times g$ for 3 min to
21 separate the two layers. The CH_2Cl_2 layer was separated, evaporated under N_2 at 40 $^\circ\text{C}$, and
22 reconstituted in 100 μL of MeCN to afford a solution of ^{18}O -labeled C-CTX1/2. Buffers of
23 various pH were assessed for their effects on label stability. Buffers (100 mM) were prepared at

1 pH 3.0 (formate), pH 5.0 (acetate), pH 6.7 (ammonium acetate), pH 7.0 (phosphate) and pH 9.0
2 (ammonium bicarbonate). ^{18}O -labeled C-CTX1/2 (20 μL) was mixed with 1:1 MeCN–buffer (90
3 μL), placed in the HPLC autosampler at 25 $^{\circ}\text{C}$, analyzed by LC–HRMS at regular intervals for
4 17 h, and the isotopic profile assessed with the NRC Isotope Enrichment Calculator to assess the
5 stability of the isotope labeling of CTXs overnight.

6 Gambierone and 44-methylgambierone (10 μL) were aliquoted together into a vial and
7 evaporated under N_2 at 35 $^{\circ}\text{C}$. The residue was dissolved in 8:4:1 MeCN– H_2^{18}O –formic acid
8 (100 μL) and allowed to stand at ambient temperature for 48 h. The solvent was evaporated
9 under N_2 and the residue dissolved in CHCl_3 (250 μL) and prepared mAPBAG was added (100
10 μL , filtered)(Mudge et al., 2022). The suspension was shaken for 3 h at 850 rpm and ambient
11 temperature. The CHCl_3 was removed with a micropipette and residual solvent was gently
12 evaporated from the gel under N_2 . The gambierones were eluted from the gel by adding 1:1
13 MeCN– H_2O (250 μL) and shaking the suspension at 850 rpm for 2 h. The resulting solution of
14 ^{18}O -labeled gambierones was transferred to a glass vial with a micropipette and stored at -20°C
15 until the stability assessment. Strongly acidic conditions (HCl, 100 mM; pH 1.0) and various pH
16 buffers (100 mM) were prepared at pH 3.0 (formate), pH 5.0 (acetate), pH 6.7 (ammonium
17 acetate), pH 7.0 (phosphate) and pH 9.0 (ammonium bicarbonate) to assess the stability of ^{18}O -
18 labeled gambierones. Aliquots of the solution of ^{18}O -labeled gambierones (10 μL) were
19 combined with 1:1 MeCN–buffer (90 μL), placed in the HPLC autosampler at 25 $^{\circ}\text{C}$, and
20 analyzed by LC–HRMS at regular intervals for 17 h. The observed isotopic patterns were then
21 evaluated with the NRC Isotope Enrichment Calculator to assess the stability of the isotope
22 labeling of gambierones.

1 2.8. Kinetic Modeling and Data Fitting

2 The changes in the isotopic composition of gambierones were modelled using a network
 3 of reversible first-order reactions. The two sites of exchangeable oxygen give rise to four distinct
 4 isotopologues depending on the incorporation of oxygen-18: unlabeled gambierone ($^{16}\text{O}-^{16}\text{O}$ or
 5 y_1), two mono-labeled gambierones ($^{16}\text{O}-^{18}\text{O}$ and $^{18}\text{O}-^{16}\text{O}$ or y_2 and y_3), and bi-labeled
 6 gambierone ($^{18}\text{O}-^{18}\text{O}$ or y_4).



7

8 Scheme 1. The kinetic model for oxygen-18 exchange in gambierones.

9

10 The concentration profiles of these four isotopologues was modelled using a set of four
 11 ordinary differential equations with three parameters (eq. 1) – forward rate constants
 12 corresponding to the uptake of oxygen-18 at each site (k_1 and k_2) and a scale parameter (R) which
 13 sets the magnitude of the two backward rate constants (k_1/R and k_2/R).

14

$$15 \begin{bmatrix} dy_1/dt \\ dy_2/dt \\ dy_3/dt \\ dy_4/dt \end{bmatrix} = \begin{bmatrix} -k_1 - k_2 & R^{-1}k_1 & R^{-1}k_2 & 0 \\ k_1 & -R^{-1}k_1 - k_2 & 0 & R^{-1}k_2 \\ k_2 & 0 & -k_1 - R^{-1}k_2 & R^{-1}k_1 \\ 0 & k_2 & k_1 & -k_1 - k_2 \end{bmatrix} \times \begin{bmatrix} y_1(t) \\ y_2(t) \\ y_3(t) \\ y_4(t) \end{bmatrix} \quad (1)$$

16

1 This mechanistic kinetic model was fitted to the parsed mass spectral data in R using a
2 general-purpose quasi-Newton optimization method. For example, the observed proportion of
3 mono-labeled gambierones at any given time is calculated from the model parameters as $[y_2(t) +$
4 $y_3(t)]/[y_1(t) + y_2(t) + y_3(t) + y_4(t)]$. The kinetic model was fitted to parsed mass spectra by finding
5 the best parameter values that minimized the squared differences between the observed and
6 predicted proportions of non-labeled, mono-labeled, and bi-labeled gambierones across all time
7 points.

8 2.9. Stability Data Fitting

9 Stability data from Section 2.6 were plotted in SigmaPlot (version 14.0) where 3-
10 parameter exponential decay curves were fitted to the stability data with constraints to the
11 variables y_0 and b , which were set to 0.2% (the natural abundance of ^{18}O), and > 0 (eq. 2),
12 respectively.

13

$$14 \text{ Extent of Labeling (\%, stability)} = y_0 + a \times e^{(-b \times t)} \quad (2)$$

15

16 Observed half-lives ($t_{1/2}$) were calculated from the first order rate constants (b) obtained from
17 fitting the stability data.

18

19 3. Results and Discussion

20 3.1. ^{18}O -Labeling of C-CTXs

21 The hemiketal on the N-ring at C-56 on C-CTX1/2 (Figure 2) opens under acidic
22 conditions, as suggested by the formation of a methyl ketal in acidified methanol (Estevez et al.,
23 2020a), and by the apparent on-column epimerization during LC–HRMS (Kryuchkov et al.,

1 2020). This potentially allows for exchange with H_2^{18}O to produce ^{18}O -labeled C-CTXs.
2 Preliminary work was performed with aliquots of fish extracts contaminated with C-CTXs and
3 monitored using LC–HRMS. The extent of labeling of C-CTX1/2 was assessed by analyzing the
4 isotope distribution of adduct ions in the full-scan mass spectra using the NRC Isotope
5 Enrichment Calculator. Full-scan mass spectra (Figure 1) obtained from *Experiment 1* revealed
6 an apparent variation in ^{18}O -labeling in the adducts, with the isotope distribution of $[[^{18}\text{O}]\text{M}+\text{H}]^+$
7 and the ammonium adduct indicating 52 % and 71 % labeling, respectively. The variance
8 between the observed extent of labeling for the $[[^{18}\text{O}]\text{M}+\text{H}]^+$ and the ammonium adduct is most
9 likely due to the facile loss of water in the electrospray ion source in positive mode. The loss of
10 H_2O or H_2^{18}O from the ammonium adduct interferes with the observed isotope distribution of the
11 $[[^{18}\text{O}]\text{M}+\text{H}]^+$ ion. The labeling percentages were essentially identical when determined from the
12 ammonium, sodium and potassium adducts (data not shown) in the full-scan spectrum, and were
13 consistently higher than those determined from the corresponding $[\text{M}+\text{H}]^+$ ions, suggesting that
14 the extent of labeling determined using the ammonium adduct should be used. The lower
15 apparent labeling of the in-source water loss fragment at m/z 1123.6234 ($[\text{M}+\text{H} - \text{H}_2\text{O}]^+$ (Figure
16 1) suggested that the labeling was at a location where it was readily lost from $[\text{M}+\text{H}]^+$ under ESI
17 conditions. Due to the relatively low abundance of the $[[^{18}\text{O}]\text{M}+\text{H}]^+$ for the labeled CTX, it was
18 not possible to obtain MS/MS spectra to confirm the location of label, but based on the presence
19 of a hemiketal at C-56, the N-ring hemiketal is the only plausible location for the addition of the
20 labeled oxygen.

21 The extent of labeling was relatively low after 48 h, therefore a partial factorial experiment
22 was conducted to determine the factors responsible for the low extent of labeling observed in the
23 preliminary experiment. These included a comparison of the proportion of ^{18}O -water, and the

1 type and concentration of the acid used. The results comparing the average response of the low
 2 and high levels for each factor, as summarized in Table 1 and Figure S1, indicated that lower
 3 percentages of labeling were observed with trifluoroacetic acid and higher acid concentration,
 4 and increased percentages with higher proportions of ^{18}O water, while temperature did not have a
 5 significant effect. Further investigations on the proportions of water were later found to have
 6 minimal effect on labeling, as it was present in excess relative to C-CTX1/2. Based on these
 7 findings, a 24-h kinetic study was performed to assess the rate of labeling (*Experiment 2*). Given
 8 the low ^{18}O -incorporation observed after 48 h and the limited supply of C-CTX1/2, these
 9 samples were placed in the autosampler at 25 °C and analyzed after 8 h, and every 4 h thereafter.
 10 At 8 h, labeling had reached 73% based on the ammonium adduct and did not change throughout
 11 the remainder of the experiment, therefore no kinetic modeling was possible for C-CTXs. These
 12 results suggested that ^{18}O -exchange with C-CTX1/2 might be fast, and that the low incorporation
 13 observed could be caused by back-exchange (loss of ^{18}O) prior to reaching the mass
 14 spectrometer. This would be consistent with the lower extent of labeling observed with TFA
 15 during the factorial study, as the stronger acid would be expected to promote faster back-
 16 exchange.

17

18 **Table 1.** Parameters evaluated and the overall level of ^{18}O incorporation into C-CTX1/2.

Trial	Acid type	Acid (%)	H_2^{18}O (%)	Temperature (°C)	Extent of labeling (%) $[\text{M}+\text{NH}_4]^+$
1	Trifluoroacetic acid	5	40	25	30.6
2	Trifluoroacetic acid	2.5	20	40	35.2
3	Formic acid	5	20	40	60.8
4	Formic acid	2.5	40	25	73.8

19

20 This hypothesis was tested by reducing the labeled C-CTX1/2 to C-CTX3/4 with sodium
 21 borohydride (Kryuchkov et al., 2020), which in the case of C-CTX1/2 labeled on the ketal at C-

1 56, will result in a non-exchangeable ^{18}O -labeled hydroxy group at C-56 (*Experiment 3*; Figure
2 2). For this work, semi-purified C-CTX1/2 without detectable levels of C-CTX3/4 was used in
3 order to avoid interference by any unlabeled C-CTX3/4 present in the sample. Comparison of the
4 full-scan mass spectra of naturally-occurring C-CTX3/4 in a fish extract and the borohydride-
5 reduced [^{18}O]C-CTX1/2 (i.e. ^{18}O -labeled C-CTX3/4) indicated that the labeling of C-CTX1/2
6 had occurred very quickly, with greater than 90% ^{18}O -labeling observed after 2 h (Figure 3).
7 Furthermore, the resulting [^{18}O]C-CTX3/4 did not undergo back-exchange prior to detection.
8 This supports the hypothesis that the measured ^{18}O -incorporation of C-CTX1/2 had been affected
9 by back-exchange promoted by the presence of acid and exposure to unlabeled water in the
10 chromatographic separation. That greater than 90% labeling was observed after 2 h of reaction
11 suggested this reaction was very fast, with a half-life of less than 30 min. Furthermore, the
12 product-ion spectrum of [^{18}O]C-CTX3/4 indicated that the location of label must be the open N-
13 ring, as shown by the presence of unlabeled product ion at m/z 979.5395 and ^{18}O -labeled product
14 ions at m/z 285.1943, 257.1530 and 227.1524 (Figure S2). Therefore, C-CTX1/2 became
15 irreversibly ^{18}O -labeled at the C-56 hydroxy group when reduced to ^{18}O -labeled C-CTX3/4
16 (Figure 3), confirming the position of the label at C-56.

17 A procedure was developed to extract the labeled C-CTX1/2 by liquid-liquid partitioning
18 (*Experiment 4*) with CH_2Cl_2 in the absence of unlabeled water, to separate it from the acid. This
19 allowed the recovery of the labeled C-CTX1/2 under neutral conditions and resulted in a 91%
20 incorporation of ^{18}O into C-CTX1/2, based on the ammonium adduct (Figure 4).

21 3.2. ^{18}O -Labeling of Gambierones

22 Gambierone and 44-methylgambierone have two locations in their structures potentially
23 available for oxygen exchange under acidic conditions. These are the hemiketal located at C-4 on

1 the A-ring and the ketone at C-40 on the aliphatic hydroxyketone side chain (Figure 5). Initial
2 investigations of both gambierones (*Experiment 1*; Figure 6) resulted in 88% and 91% labeling at
3 two positions in the molecules.

4 The product-ion spectrum of ^{18}O -labeled gambierone had several product ions in the high
5 mass range that were 2 or 4 m/z higher compared to unlabeled gambierone, although the MS/MS
6 data suggested that one of the labeled oxygen atoms was eliminated from the structure (Figure
7 7). There were limited product ions indicative of cleavages in the A–E rings, making
8 identification of the exact location of this easily eliminated label difficult. However, based on the
9 structure of gambierone, it is most likely the A-ring hemiketal at C-4. The low-mass product ions
10 at m/z 221.1417, m/z 291.1832 and m/z 345.2301 strongly suggested labeling on or after the I-
11 ring, with the most probable location being the ketone at C-40. The product ion at m/z 161.0960
12 resulting from the cleavage between C-36 and C-37 in the I-ring was present in both the labeled
13 and unlabeled spectra, suggesting that the water loss for this product ion occurred from the
14 ketone position. Corresponding product ions were observed in the mass spectra of $[^{18}\text{O}_2]$ 44-
15 methylgambierone and unlabeled 44-methylgambierone, suggesting that the locations of the ^{18}O -
16 labels in 44-methylgambierone were identical to those in gambierone (Figure S3).

17 A kinetic study (*Experiment 2*) was performed to assess the rate of labeling for
18 gambierones at 25 °C. After 13 h, the extent of labeling reached a maximum, with 96% and 89%
19 labeling at locations 1 and 2, respectively. This corresponded to approximately 13% labeling at a
20 single location, and 83% at both locations, in the two gambierones at the completion of the
21 experiment (Figure 8; Figure S4). The labeling reactions followed first-order kinetics and were
22 fitted to the network of reversible first-order reactions, with logarithmic decay and growth
23 curves. The kinetics of isotope exchange were virtually identical for both compounds, with

1 unlabeled gambierone and 44-methylgambierone incorporating oxygen-18 with half-lives of
2 about 40 min, and with the two exchange sites being labeled with half-lives of approximately 50
3 and 190 min (Figures 8 and S4). An additional kinetic study of 44-methylgambierone was
4 monitored over a 2-h period to identify which position was exchanging at a faster rate by
5 acquiring MS/MS data of the labeled products. Monitoring the product-ion spectrum of
6 $[[^{18}\text{O}]\text{M}+\text{H}]^+$ at m/z 1041.4973 and the rate of formation of the product ion at m/z 235.1579
7 indicated that the ketone at C-40 exchanged faster than the hemiketal at C-4.

8 Sodium borohydride reduction was used to test for the possible effects of back-exchange
9 during analysis due to the presence of acid, as was done for C-CTX1/2 (*Experiment 3*). The
10 unreduced ^{18}O -labeled gambierones showed 94% labeling at both locations. Gambierones
11 contain two functional groups that are potentially reducible with NaBH_4 , the carbonyl at C-40
12 and the hemiketal at C-4, and these are also the expected sites for oxygen-18 labeling.
13 Borohydride-reduced ^{18}O -labeled gambierones were observed to have 94% labeling at both
14 locations (Figure 9), suggesting that no detectable back-exchange had occurred prior to
15 detection, in contrast to the situation for C-CTX1/2.

16 3.3. Isotopic Stability of ^{18}O -labeled C-CTXs and Gambierones

17 Due to the limited quantities of semi-purified C-CTX1/2 available, the stability of the ^{18}O -
18 labeling could only be followed at a few pH values and analyzed 3 to 4 times over a 17 h period
19 at 25 °C. The labeled C-CTX1/2 used in the stability study had 78% labeling at a single location
20 for C-CTX1/2. LC–HRMS analysis indicated an incorporation level of 78% for the ^{18}O -labeled
21 C-CTX1/2 after 17 h at pH 6.7, indicating very high stability under neutral conditions, whereas
22 all other pH conditions assessed resulted in some degree of back-exchange (Table 2). The fastest
23 back-exchange was observed at pH 3 and 9, with ^{18}O -labeled C-CTX1/2 having half-lives of less

1 than 24 h. Full-scan HRMS spectra at each pH after 17 h are shown in Figure S5, and confirm
 2 considerable back-exchange under these conditions. Neutral conditions were also evaluated with
 3 phosphate buffer (100 mM). These data suggest that there may be some phosphate-catalyzed
 4 back-exchange of [¹⁸O]C-CTX1/2, as after 17 h the extent of labeling dropped to 63% in the
 5 presence of phosphate buffer, which was prepared at a neutral pH similar to that of the
 6 ammonium acetate for which minimal back-exchange was observed. Due to limited quantities of
 7 C-CTX1/2, lower concentrations of phosphate buffer were not assessed. However, a previous
 8 study reported concentration-dependent general acid catalysis of the exchange of the carbonyl
 9 oxygen of acetone (Greenzaid et al., 1968). Based on these findings, ¹⁸O-labeled C-CTX1/2
 10 appears to be sufficiently stable under neutral conditions to be used for analytical measurements
 11 and possibly also for in vitro assays, although alternative neutral buffers may be necessary if the
 12 use of phosphate buffers proves problematic. While not assessed due to sample availability, it is
 13 probable that storage at lower temperatures would reduce the rate of back-exchange.

14 **Table 2.** Stability of [¹⁸O]C-CTX1/2 under several pH conditions, based on LC–HRMS
 15 measurement of the ammonium adduct [M+NH₄]⁺, at the beginning and completion of a stability
 16 study (17 h) at 25 °C, with observed half-life estimates based on 3-parameter logarithmic decay
 17 curves.*

pH	buffer	Extent of labeling (%)		Half-life (h)
		initial	after 17 h	
3.0	formate	77.9	20.5	11
5.0	acetate	78.9	75.2	220
6.7	ammonium acetate	78.5	78.6	3900
7.0	phosphate	76.8	62.9	55
9.0	ammonium bicarbonate	77.8	20.6	11

18 *constrained to $y_0=0.2\%$

19 To evaluate the rate of back-exchange for ¹⁸O-labeled C-CTX3/4, [¹⁸O]C-CTX1/2 was
 20 reduced with sodium borohydride and mixed with formate (pH 3), ammonium acetate (pH 6.7)
 21 and ammonium bicarbonate (pH 9) buffers. The samples were analyzed after 5 h at ambient
 22 temperature and all were found to have >95% ¹⁸O-label incorporation, showing that back-

1 exchange was negligible, in contrast to [^{18}O]C-CTX1/2. This is because borohydride reduces the
2 hemiketal at C-56 of C-CTX1/2 to an open-ring hydroxy group (Kryuchkov et al., 2020), where
3 the ^{18}O -label is permanently affixed to the molecule and no longer able to undergo acid-
4 catalyzed oxygen exchange.

5 As with C-CTXs, the stability of gambierones was assessed after the removal of residual
6 acid. However, the procedure developed for separation of the C-CTXs from the acid catalyst was
7 not appropriate for gambierones, because they do not partition efficiently into CH_2Cl_2 (Estevez et
8 al., 2020b). Instead, the labeled gambierone was removed from the acidic solution using a
9 recently developed boronate affinity technique with mAPBAG (Mudge et al., 2022). The
10 resulting solution was a mixture of the two ^{18}O -labeled gambierones in 1:1 MeCN–water,
11 thereby enabling the addition of buffered solutions to control the pH. The stability of the ^{18}O -
12 incorporation was assessed at a range of pH values, from strongly acidic to weakly basic for
13 gambierone and 44-methylgambierone. Comparisons of the full-scan spectra of the time zero
14 control with spectra after 17 h in the various buffers are shown in Figures S6 and S7. The extent
15 of labeling at the two positions was determined from the LC–HRMS spectra with the NRC
16 Isotope Enrichment Calculator and fitted to 3-parameter first-order decay curves at each pH
17 (Figures S8 and S9), and the results are summarized in Table 3. Back-exchange (loss of ^{18}O) was
18 observed relatively quickly in strong acid and at pH 3 and 9, indicating that these are not suitable
19 for storage or use of ^{18}O -labeled gambierones. Strong acid (0.1 M HCl) caused a rapid back-
20 exchange at both locations on the gambierone structure, with the ^{18}O at C-40 exchanging almost
21 instantaneously, with a half-life of around 1 min, while for the C-4 label the half-life was 22 min.
22 This back-exchange was slower at pH 3 and 9, but loss of labeling was also observed. As was
23 observed for C-CTXs, phosphate buffer appeared to have a catalytic effect on the stability of the

1 label. Exchange was slower at pH 5 with acetate buffer (half-life ~550 h) than at pH 7 using
 2 phosphate (~ 175 h). The half-life estimated for ¹⁸O-labeled gambierones in ammonium acetate
 3 at neutral pH was approximately 100–130 d. These experiments were performed at 25 °C and,
 4 although it is likely that stability would be improved at lower temperatures, the stability of the
 5 label may be sufficient for analytical measurements and in vitro assays.

6
 7 **Table 3.** Effect of pH on ¹⁸O-labeled gambierone and 44-methylgambierone in a 17-h stability
 8 study at 25 °C, and observed *t*_{1/2} for back-exchange of the two labeled positions (C-4 and C-40).

pH	Gambierone						44-Methylgambierone					
	Half-life (h)		Extent of Labeling (%)				Half-life (h)		Extent of Labeling (%)			
	C-40	C-4	Initial		17 h		C-40	C-4	Initial		17 h	
			C-40	C-4	C-40	C-4			C-40	C-4	C-40	C-4
1.0 [†]	0.4	0.02	91	91	15 [†]	0 [†]	0.4	0.02	91	91	16 [†]	0 [†]
3.0	260	9	91	91	87	17	270	8	91	91	87	16
5.0*	570	570	91	91	88	88	540	540	90	90	88	88
6.7*	3100	3100	91	91	90	90	2400	2400	91	91	90	90
7.0*	170	170	91	91	84	84	180	180	90	90	84	84
9.0	45	13	90	90	67	29	44	13	90	90	68	28

9 [†]Initial and final (45 min after preparation) analysis for pH 1.

10 *Slow back-exchange appeared consistent at both positions, but was too slow to measure
 11 accurately at these pH values in only 17 h.

12
 13 There remains a paucity of CTX reference materials, which are necessary for reliable
 14 quantitation, identification and verification of these toxins in screening and monitoring work, and
 15 in fish from outbreaks worldwide. Recent work on the development of CTX reference materials
 16 has highlighted several difficulties associated with this work, including large sample
 17 requirements, low levels of CTXs in the fish flesh, and relatively low LC–MS instrument
 18 response (Gago-Martinez et al., 2021). Isotope-labeled internal standards provide a
 19 complementary approach for the development of sample preparation and quantitative procedures,
 20 as isotope dilution is an effective methodology for evaluating matrix effects, extraction

1 efficiencies, and instrument response in high matrix materials (Haddad et al., 2019; Stokvis et
2 al., 2005). This would require less standard for the developmental stages of LC–MS methods,
3 thus reducing overall reference material needs. Future work will focus on using ¹⁸O-labeled
4 gambierones and CTXs to establish isotope dilution methodologies and determine the impact of
5 matrix effects on LC–MS detection of these toxins in crude and semi-purified fractions from
6 algae and fish.

7

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21

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1 List of Figure Captions

2 **Figure 1.** Full-scan mass spectra in positive ionization mode for C-CTX1/2 in: (A) unlabeled
3 control, and; (B) ^{18}O -labeled C-CTX1/2 after 48 h of reaction (*Experiment 1*), using an extract of
4 *S. barracuda* containing C-CTX1/2.

5 **Figure 2.** Sodium borohydride reduction of ^{18}O -labeled C-CTX1/2 to produce labeled C-
6 CTX3/4.

7 **Figure 3.** Full-scan mass spectra in positive ionization mode for: (A) unlabeled C-CTX3/4, and;
8 (B) ^{18}O -labeled C-CTX3/4 produced by sodium borohydride reduction of ^{18}O -labeled C-CTX1/2
9 (*Experiment 3*).

10 **Figure 4.** Full-scan mass spectra in positive ionization mode for: (A) unlabeled C-CTX1/2, and;
11 (B) ^{18}O -labeled C-CTX1/2 after CH_2Cl_2 partitioning to remove residual acid (*Experiment 4*).

12 **Figure 5.** Chemical structures of gambierone and 44-methylgambierone.

13 **Figure 6.** Full-scan mass spectra in negative ionization mode for: unlabeled (A) gambierone and
14 (B) 44-methylgambierone; and ^{18}O -labeled (C) gambierone and (D) ^{18}O -labeled 44-
15 methylgambierone after 48 h of acid catalyzed reaction with H_2^{18}O (*Experiment 1*).

16 **Figure 7.** Product-ion spectra in positive ionization mode of the protonated molecules of: (A)
17 gambierone ($[\text{M}+\text{H}]^+$, m/z 1025.4750), and; (B) ^{18}O -labeled gambierone ($[\text{M}+\text{H}]^+$, m/z
18 1029.4844).

19 **Figure 8.** ^{18}O -incorporation into gambierone with time. (A) Extent of labeling at each location,
20 and; (B) proportion of gambierone molecules labeled at no, one, or both locations within the
21 molecule (*Experiment 2*). The decay and growth curves were fitted to the data using a network of
22 reversible first-order reactions (Scheme 1).

23 **Figure 9.** Full-scan mass spectra in negative ionization mode for: (A) ^{18}O -labeled gambierone,
24 and; (B) 44-methylgambierone, and after sodium borohydride reduction of; (C) ^{18}O -labeled
25 gambierone, and; (D) 44-methylgambierone, to irreversibly incorporate stable isotopes.

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