Isolation of Alpha-Toxin-Deficient Clostridium perfringens Type F from Sewage Influent and Effluents

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ABSTRACT Clostridium perfringens is classified into types A to G, and all types produce alpha-toxins; however, C. perfringens type F that is negative for phospholipase C (PLC) activity of alpha-toxin has been isolated from the environment and cases of humans afflicted by food poisoning. This study aimed to elucidate the distribution of PLC-negative C. perfringens type F in sewage influents and effluents. Influent and effluents of two wastewater treatment plants were collected monthly between July 2016 and January 2020 and between August 2018 and January 2020, respectively. Isolation rates of PLC-negative C. perfringens type F from sewage influents and effluents were 38% (33/86) and 22% (8/36), and the numbers of isolates were 43 and 13, respectively. The locus of the enterotoxin gene of all isolates was determined to be in a plasmid with an IS1151 sequence, and multilocus sequence typing revealed that all 17 representative isolates were assigned as sequence type 186. Sequencing of the plc gene of these representative isolates showed that nonsense mutation (p.W98*) causing alpha-toxin deficiency should be responsible for a loss of PLC enzymatic activity. These results suggest that alpha toxin-deficient C. perfringens type F is distributed in living and water environments since sewage influents contain community wastewater, and effluents contaminate the environment. Detection of C. perfringens type F, independent of PLC activity, should be carried out on human and environmental samples.

IMPORTANCE Understanding the diversity of biochemical characteristics that may affect the identification of bacteria is essential. C. perfringens is a ubiquitous bacterium found in the environment, humans, and animals and is responsible for infectious disease in the intestine. Although the alpha-toxin of C. perfringens may be used for its detection, variants of the alpha-toxin lacking its activity have been isolated from soil and humans experiencing symptoms of diarrhea. It is valuable to disclose the prevalence of the alpha-toxin variant in the sewage of wastewater treatment plants, as it may reflect the hygienic condition of the community, as it would be a pollution source for the environment. This study shows the persistent existence and genetic characteristics of the alpha-toxin variant in sewage and reveals a lacking mechanism of the alpha-toxin activity and proposes the detection method of C. perfringens, independent of the alpha-toxin activity.

KEYWORDS alpha-toxin, phospholipase C negative, lecithinase negative, wastewater treatment plant, multilocus sequence typing

C. perfringens is an anaerobic, spore-forming bacterium that is classified as type A to G based on toxin production, including alpha, beta, epsilon, iota, the C. perfringens enterotoxin (CPE), and NetB toxin production (1). C. perfringens type F produces an alpha-toxin and CPE and is responsible for foodborne and nonfoodborne infection (2, 3). Although all types of C. perfringens produce an alpha-toxin which exhibits phospholipase C (PLC) and sphingomyelinase activity (4), C. perfringens that
TABLE 1 Characteristics of PLC-negative C. perfringens type F as assessed by MLST

| Strain | Location of cpe | cpb2 | Source | WWTP | Mo and yr of isolation | ST |
|--------|----------------|------|--------|------|------------------------|----|
| 17-97  | Plasmid with an IS1151 sequence | +    | Influent A | April 2017 | 186 |
| 17-113 | Plasmid with an IS1151 sequence | +    | Influent B | April 2017 | 186 |
| 18-67  | Plasmid with an IS1151 sequence | +    | Influent A | December 2017 | 186 |
| 18-104 | Plasmid with an IS1151 sequence | +    | Influent B | January 2018 | 186 |
| 18-108 | Plasmid with an IS1151 sequence | −    | Influent B | February 2018 | 186 |
| 18-266 | Plasmid with an IS1151 sequence | +    | Influent A | August 2018 | 186 |
| 18-331 | Plasmid with an IS1151 sequence | +    | Influent A | October 2018 | 186 |
| 19-37  | Plasmid with an IS1151 sequence | +    | Influent A | February 2019 | 186 |
| 19-41  | Plasmid with an IS1151 sequence | −    | Influent B | February 2019 | 186 |
| 19-98  | Plasmid with an IS1151 sequence | +    | Influent B | April 2019 | 186 |
| 19-198 | Plasmid with an IS1151 sequence | +    | Influent A | November 2019 | 186 |
| 19-195 | Plasmid with an IS1151 sequence | +    | Influent B | November 2019 | 186 |
| 18-280 | Plasmid with an IS1151 sequence | +    | Effluent A | October 2018 | 186 |
| 18-283 | Plasmid with an IS1151 sequence | +    | Effluent B | October 2018 | 186 |
| 18-348 | Plasmid with an IS1151 sequence | +    | Effluent A | January 2019 | 186 |
| 19-61  | Plasmid with an IS1151 sequence | +    | Effluent B | April 2019 | 186 |
| 19-203 | Plasmid with an IS1151 sequence | +    | Effluent B | December 2019 | 186 |

* +, positive; −, negative.

produces an alpha-toxin negative for PLC activity and is a plc-gene variant was isolated from Antarctic soil (5). In addition, previous studies confirmed isolations of PLC-negative C. perfringens type F from food poisoning patients (6, 7). However, these isolates are not typical among human isolates because the detection of PLC activity with egg yolk agar may be used to detect C. perfringens (5, 6, 8); thus, little is known about the distribution of PLC-negative C. perfringens type F among humans and other sources, such as the environment.

In a previous study, Salmonella spp. isolated from sewage influents of wastewater treatment plants (WWTPs) were more variable than those isolated from humans (9). Additionally, genetically variable isolates of C. perfringens type F were recovered from sewage influents and effluents of WWTPs (10, 11). Therefore, sewage may contain variable C. perfringens type F, such as PLC-negative strains. Furthermore, the distribution of PLC-negative strains may be estimated based on sewage samples, which would contain community wastewater and contaminate the water environment (11).

The aim of this study was to reveal the isolation rate, genetic characteristics, and plc sequence of PLC-negative C. perfringens type F isolates from sewage influents and effluents of WWTPs to reveal the distribution of these C. perfringens strains in the water environment and to detect the PLC-negative mechanism.

RESULTS

Isolation of PLC-negative C. perfringens type F. The isolation rates of PLC-negative C. perfringens type F from sewage influents and effluents of two WWTPs (WWTP-A and WWTP-B) in Yamanashi, Japan, were 38% (33/86) and 22% (8/36), and the number of isolates from these was 43 and 13, respectively (Table S1 in the supplemental material). All isolates except for two isolates from two sewage influents of WWTP-B were positive for the C. perfringens beta2 toxin gene (cpb2). There was no significant difference between WWTP-A and WWTP-B in the isolation rate and number of isolates (P > 0.05), and no seasonal trend was observed.

Characterization of the isolates. Characterization of the isolates was conducted by cpe genotyping to determine the locus of cpe and multilocus sequence typing (MLST). The locus of cpe of all isolates was discovered in a plasmid with an IS1151 sequence downstream of cpe. MLST demonstrated that all representative 17 isolates were assigned as sequence type (ST) 186 (Table 1).

Analysis of the plc sequence. Analysis of the plc sequence was performed to determine the PLC-negative mechanism and alpha-toxin sequence type. The sequencing of plc revealed that all isolates were identical to each other and showed a nucleotide
substitution at nucleotide position 293 (c.293G>A) compared with the plc sequence of CP228 (12) (Fig. S1). This nucleotide substitution resulted in a nonsense mutation (p.W98*) in the deduced amino acid sequence, which was determined to be a single amino acid substitution of the alpha-toxin sequence type Ve and was designated alpha-toxin sequence type N (nonsense mutation) (Table 2 and Fig. S2).

DISCUSSION

In the present study, we revealed the presence and genetic characteristics of PLC-negative C. perfringens type F isolates from sewage influents and effluents of WWTPs and elucidated their PLC-negative mechanism. Although PLC-negative C. perfringens has rarely been reported since PLC activity is an important criterion for the identification of C. perfringens (5, 6, 8), the present study showed that PLC-negative C. perfringens type F was isolated from 38% and 22% of sewage influents and effluents, respectively. Our previous study revealed that the isolation rates of PLC-positive C. perfringens type F from sewage influents and effluents were 80% (69/86) and 56% (20/36), respectively (11), which was higher than that of PLC-negative C. perfringens type F observed in the present study. These results suggest that even though PLC-negative isolates occur less among C. perfringens type F strains, the presence of PLC-negative isolates in water environments such as sewage should be considered.

The plc sequencing of all isolates showed a nucleotide substitution (c.293G>A) compared with the plc sequence of CP228 (12) (Fig. S1). This nucleotide substitution resulted in a nonsense mutation (p.W98*) in the deduced amino acid sequence, which was determined to be a single amino acid substitution of the alpha-toxin sequence type Ve and was designated alpha-toxin sequence type N (nonsense mutation) (Table 2 and Fig. S2).

### TABLE 2 Alpha-toxin sequence types and their amino acid positions that were different from those of strain 13

| Alpha-toxin sequence type | Amino acid at position: a |
|---------------------------|---------------------------|
|                           | 13 | 15 | 22 | 47 | 54 | 71 | 98 | 149 | 195 | 202 | 205 | 365 | 373 |
| Strain 13                 | T  | A  | A  | V  | L  | E  | W  | L  | A  | D  | A  | A  | I  |
| Type I                    | A  |    |    |    |    |    |    |    |    |    |    |    |    |
| Type II                   | A  |    |    |    |    |    |    |    |    |    |    |    | V  |
| Type III                  | V  | V  | M  | D  | I  |    |    |    |    |    |    |    | V  |
| Type IV                   | A  |    |    |    |    |    |    |    |    |    |    | M  | V  |
| Type V                    | A  |    |    |    |    |    |    |    |    |    |    | A  | T  |
| Type Ve                   | A  |    |    |    |    |    |    |    |    |    |    | A  | T  |
| Type VI                   |    | V  | N  | V  |    |    |    |    |    |    |    |    |    |
| Type VII                  | A  | S  | I  |    | V  |    |    |    |    |    |    |    |    |
| Type VIII                 |    | V  |    |    |    |    |    |    |    |    |    |    |    |
| Type IX                   | I  |    |    |    |    |    |    |    |    |    |    |    |    |
| Type X                    | I  |    |    |    |    |    |    |    |    |    |    |    |    |
| Type Nc                   | A  |    |    |    |    |    |    |    |    |    |    |    |    |

a Alpha-toxin sequence type strain 13 and types I to V, type Ve, and types VI to X were defined by Sheedy et al. (19), Matsuda et al. (12), and Ablidgaard et al. (20), respectively.
b Amino acid positions different from those of strain 13.
c Type N (nonsense mutation) as defined in the present study.
d Amino acid in parentheses indicates the deduced amino acid if there was not a stop codon at position 98.
e Asterisk shows a stop codon.

According to the table, the nucleotide substitution at nucleotide position 293 (c.293G>A) compared with the plc sequence of CP228 (12) (Fig. S1). This nucleotide substitution resulted in a nonsense mutation (p.W98*) in the deduced amino acid sequence, which was determined to be a single amino acid substitution of the alpha-toxin sequence type Ve and was designated alpha-toxin sequence type N (nonsense mutation) (Table 2 and Fig. S2).
sequence type N is most prevalent among alpha-toxin-deficient isolates in Japan due to these results and the isolation rate of isolates in the present study, more investigations about the pathogen are necessary to discuss the situation.

Although representative isolates were examined by MLST, the results of the cpe genotyping assay and MLST indicated that the locus of cpe and the ST of the isolates in the present study were identical (a plasmid with IS\text{1151} sequence and ST186, respectively), which indicates that strains with common characteristics are widely distributed in water environments. When this is not the case, these strains show higher resistance than other alpha-toxin-deficient \textit{C. perfringens} type F against environmental stress in sewage pipes and WWTPs, even though it is not identified whether a nonsense mutation of the isolates is related to this resistance. We believe that both hypotheses occur simultaneously since it is unlikely that these strains with the same sequence type (ST) flowed into two sewage pipes independently without wide distribution and were not detected from sewage effluents, which were processed with sewage treatments, without higher stress resistance.

When alpha-toxin-deficient \textit{C. perfringens} type F isolates from sewage are derived from the community, it is likely that a substantial amount of alpha-toxin-deficient \textit{C. perfringens} type F blends into living and water environments. This is supported by the fact that \textit{C. perfringens} type F isolates from sewage influents and effluents were genetically related to those from humans and retailed bivalves (11), which accumulate \textit{C. perfringens} in the water environment (13). In addition to this, the locus of cpe of all isolates in the present study was a plasmid with an IS\text{1151} sequence, and Kiu et al. reported that strains with plasmidal cpe, including a plasmid with an IS\text{1151} sequence, were the predominant cause of food poisoning cases by \textit{C. perfringens} (3). Thus, it should be noted that the detection method of \textit{C. perfringens} type F regardless of PLC activity could be necessary for samples collected from not only the environment but also humans with gastroenteritis because PLC-negative \textit{C. perfringens} type F was isolated from food poisoning patients (6, 7), and the isolates in the present study may possess a potential for association with gastroenteritis. Future investigations that disclose the distribution of alpha-toxin-deficient \textit{C. perfringens} type F in other regions would indicate the nature of this pathogen.

A multiplex PCR for the detection of toxin genes revealed that all isolates in our study were positive for the plc gene; however, PLC production of all isolates on CW agar plates was negative. This contradiction can be explained by the single-nucleotide substitution of plc, which caused the alpha-toxin deficiency and keeps its sensitivity to primers, which enables plc detection. This result reminds us that a plc-positive strain does not mean that it is a PLC-producing strain.

The limitation of our study is that the genetic comparison of the isolates was conducted by only MLST. Whole-genome sequencing of the isolates would be required for the strict genetic comparison. Additionally, no isolate from human was analyzed in the present study. Future analyses will be essential to show the actual situation in humans.

In conclusion, alpha-toxin-deficient \textit{C. perfringens} type F was isolated from 38% and 22% of sewage influents and effluents, respectively, and the results of the MLST and cpe genotyping assay of all tested isolates were identical. Additionally, plc sequencing revealed a nonsense mutation that was estimated to be responsible for the alpha-toxin deficiency. These results suggest that alpha-toxin-deficient \textit{C. perfringens} type F is distributed in living and water environments, and the detection of \textit{C. perfringens} type F with or without PLC activity should be conducted on human and environmental samples.

**MATERIALS AND METHODS**

**Sample collection.** Sewage influents and effluents of WWTP-A and WWTP-B, which serve a population of $350,000$ and the total loads of $180,000$ m$^3$/day of wastewater in Yamanashi, Japan, were collected monthly between July 2016 and January 2020 and between August 2018 and January 2020, respectively. The collected samples were concentrated by centrifugation, as described previously (11).

**Isolation methods for PLC-negative \textit{C. perfringens} type F.** Concentrated samples were cultured in enrichment broth, as described previously (11). Colonies without PLC production were observed on a
CW agar plate (Nissui Pharmaceutical, Tokyo, Japan) that contained 50% egg yolk-enriched saline (Kyokuto, Tokyo, Japan) and were isolated and suspended in sterilized distilled water. Then, DNA was extracted by heating at 100°C for 10 min, and PCR with species-specific primers based on the 16S rRNA gene of \textit{C. perfringens} was conducted for identification of the isolates as described by Kikuchi et al. (14).

Detection of toxin genes, including \textit{C. perfringens} alpha, beta, epsilon, iota, and \textit{cpb2} and \textit{cpe}, was performed as described by van Asten et al. (15) (Table 3).

Characterization of the isolates. All isolates were characterized by \textit{cpe} genotyping assay (16) to determine the locus of \textit{cpe}, and 17 representative isolates considering month and year of isolation and source were analyzed by MLST, as described by Deguchi et al. (17) (Table 3). ST of the isolates was determined according to Xiao’s scheme (18) by submitting the sequence data to PubMLST (https://pubmlst.org/organisms/clostridium-perfringens).

Analysis of \textit{plc} sequence. To determine the PLC-negative mechanism and the alpha-toxin sequence type, \textit{plc} sequencing of 17 representative isolates was performed. Briefly, DNA was extracted by heating at 100°C for 10 min, and PCR was conducted using Thermal Cycler Dice Touch TP350 (TaKaRa Bio, Kusatsu, Japan). The amplification of \textit{plc} was performed with primer pairs reported by Sheedy et al. (19) (Table 3). Each 25-μl reaction mixture contained 2.5 μl of 10× Ex Taq buffer, 2 μl of deoxyribonucleoside triphosphate (dNTP) mixture, 0.125 μl of TaKaRa Ex Taq (TaKaRa Bio), 2 μl each of 2.5-pmol/μl primer, and 2.5 μl of template DNA. PCR products were sequenced with BigDye Terminator v3.1 cycle sequencing kit ( Applied Biosystems, Foster City, CA) according to the manufacturer’s instructions. The deduced amino acid sequence of alpha-toxin was obtained using a genetic information processing software (GENETYX ver. 13, Genetyx, Tokyo, Japan), and alpha-toxin sequence type of each isolate was assigned according to previous studies (12, 19, 20).

### Table 3: Primers used in this study

| Primer          | Sequence (5′–3′)                  | Application                      | Reference |
|-----------------|-----------------------------------|----------------------------------|-----------|
| CIPER-F         | AGATGGCCCATATCATCCAC              | Identification of \textit{C. perfringens} | 14        |
| CIPER-R         | GCAGGAGATGTCGTAATG                |                                   |           |
| CPAalphaF       | GCTAAGTTACTGGCCTGGA              | Toxinotyping                     | 15        |
| CPAalphaR       | CCTCTGATACATGCTGTAAG             |                                   |           |
| CPBetaF3        | GCCGAATCTGTAATCATCTA             |                                   |           |
| CPBetaR3        | GCGAGAAGATTGATGATATTCT           |                                   |           |
| CPBeta2totalF2  | AAATATGCTCTAACAAC                 |                                   |           |
| CPBeta2totalR   | CCAAATACTTATTATGTGAG             |                                   |           |
| CPEpsilonF      | TGGGAACTTCGATCAGG                |                                   |           |
| CPEpsilonR2     | AACTGCACTATAATTCCCTTTTCC         |                                   |           |
| CPlotaF         | AATGTTGATTATCTGCT                |                                   |           |
| CPloterR        | TTAGCAGATGCTACTCATATT            |                                   |           |
| Cpe4F           | TTCGAACTGATGTAATT                 | \textit{cpe} genotyping          | 16        |
| IS1470R1.3      | CTTCTGTGATACAGG                  |                                   |           |
| IS1470-likeR1.6 | CTTTGTGATACAGGG                  |                                   |           |
| IS1151R0.8      | ATCAAAAAATATGTTCAAGTCT           |                                   |           |
| 3F              | GATGAAAAGAGATGATGAG              |                                   |           |
| 4R              | GATGAAAAGAGATGATGAG              |                                   |           |
| gvrB-F          | ATTTGTTGACAAGATATGGAG             | MLST                             | 17        |
| gvrB-R          | ATTTCATATATNAGTTTGCC             |                                   |           |
| sigk-F          | CAATCTCTTAAAGATATGGTAATGGAG      |                                   |           |
| sigk-R          | CAGATACATATGCTTCTGATACC          |                                   |           |
| sod-F           | CAAATGATGCTACATTGTACC            |                                   |           |
| sod-R           | TTATCTATGTTAATATCTATGCC          |                                   |           |
| groEL-F         | TACAAAGATATACATTACTCTGGG          |                                   |           |
| groEL-R         | CATTCTCTTCTGGGAGATATCC           |                                   |           |
| pgk-F           | GACCTTTAGCTTCCTTAAAGATGG          |                                   |           |
| pgk-R           | CTAATCCGGATATCCTGCAGG            |                                   |           |
| nadA-F          | ATTACGACATTATATCAAATTCTCG        |                                   |           |
| nadA-R          | TTAATGCTCTTAATCTAACTC            |                                   |           |
| colA-F          | ATTAGAAATTTGTTATCAAGG             |                                   |           |
| colA-R2         | AAGACATCTATATTCTTATCGAAGC         |                                   |           |
| plc-F           | AGGAACCTATCGATGATGTAACTCT        | \textit{plc} sequencing         | 19        |
| plc-R           | GGATCATTACCTCGATACGTCG           |                                   |           |
| ss2             | CTTGAAAGAAAATCAAGG               |                                   |           |
| cpaR            | TCCTGATACATGCTGAAG               |                                   |           |
| cpaF            | GCTAATGTTACGGCGGTTGAC            |                                   |           |
| ss3             | TGTAATACCAACCAAAACC              |                                   |           |
**Statistical analysis.** The differences between WWTP-A and WWTP-B in the isolation rate and number of isolates were compared using the chi-square test, and a P value of <0.05 was considered statistically significant.

**Data availability.** The sequence data obtained in this study were deposited in the DNA Data Bank of Japan and GenBank under accession numbers LC603848 to LC604000.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**SUPPLEMENTAL FILE 1, PDF file, 0.2 MB.**

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