



# Enhanced Surveillance of *Clostridium difficile* Infection: Ireland – Q2 2018 National Report

# **Executive Summary**

- During Q2 2018, a total of 527 cases of C. difficile infection (CDI) were reported to enhanced surveillance from 55 acute public and private hospitals across Ireland.
- The national overall rate of CDI in hospitalised patients was 4.0 cases per 10,000 bed days used (BDU) [402 cases], higher than that reported for Q2 2017 [359 cases; rate = 3.7]
- There were 250 cases of CDI deemed to be hospital-acquired (HA-CDI), of which 221 were new, representing a national new HA-CDI rate of 2.2 [median rate = 1.4]
- All hospitals reported using a C. difficile testing method recommended in the 2014 updated national clinical guidelines for C. difficile
- Ribotyping data was available for 20% of cases, with ribotypes 002, 078 and 014 the most frequently reported
- With regard to acquisition, C. difficile was mostly associated with acute hospitals (250; 47%).
   However, many cases were associated with long term care facilities (LTCF) (39; 7%) and the community (134; 25%)
- CDI symptom onset occurred in the community for 41% of all cases (216):
  - o This emphasises the importance of considering CDI when evaluating any patient with potentially infectious diarrhoea in all healthcare settings, including hospitals, primary care and LTCF. Guidance on CDI for primary and long-term care settings is available at the following link:

### http://www.hpsc.ie/A-Z/Gastroenteric/Clostridiumdifficile/Guidelines/File,14387,en.pdf

- o It also emphasises the importance for all microbiology laboratories in Ireland to implement the recommendations of the national *C. difficile* clinical guidelines to routinely include *C. difficile* testing for all faeces specimens that take the shape of the container submitted from patients aged 2 years, regardless of patient location or clinician request. Guidance on *C. difficile* testing is available in Section 2.5, pages 43 54 of the national *C. difficile* clinical guidelines
- The excellent participation in enhanced surveillance since its launch in 2009 indicates the commitment of the microbiology laboratories, multi-disciplinary infection prevention and control and antimicrobial stewardship teams, along with hospital management to understanding the epidemiology of this important infection and minimising the risk of patients acquiring CDI as an unintended consequence of healthcare

# Part 1: National CDI Epidemiology – Q2 2018

CDI data was reported to the enhanced surveillance programme from 55 acute public and private hospitals across Ireland (*Appendix A*). There were 527 reported CDI cases in patients aged 2 years, of those 402 were reported in hospitalised patients, giving a national CDI rate in hospitalised patients of 4.0 cases per 10,000 bed days used (BDU), which is higher than that reported for Q2 2017 [359 cases; rate = 3.7]. The majority were aged 65 years (69%) and were female (56%). Twelve cases of severe CDI were reported (2.3%), defined as requiring critical care admission or colectomy due to complications of CDI, which comparable with twelve cases (2.7%) for Q2 2017. *Table 1* displays the breakdown of all CDI cases for Q2 2018 versus Q2 2017, by case type, origin, onset and severity. CDI case definitions are summarised in Appendix B.

## **CDI Case Type**

The majority were categorised as new infections (85%), with 9% recurrent and for 6%, the CDI case type was unknown.

## **CDI Origin**

The majority were categorised as healthcare-associated (HCA) CDI [n=308; 58%], with community-associated (CA) CDI accounting for 25% [n=134]. For the remainder, the origin either could not be determined [n=46; 9%] or was unknown [n=39; 7%]. Of 308 HCA-CDI cases, the origin was the reporting hospital, termed hospital-acquired (HA) for 250 (81%), a LTCF for 39 (13%) and 'other' or 'other healthcare facility' for 17 (6%).

#### **CDI Onset**

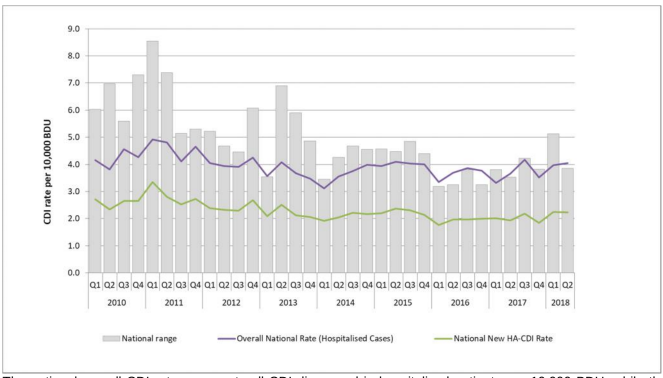
Patient locations at onset of CDI symptoms included; while admitted to a healthcare facility, termed healthcare-onset (HO) for 286 cases (54%), while residing in the community, termed community onset (CO) for 216 cases (41%) and unknown patient location for 25 cases (5%). Of 286 HO CDI cases, the reporting hospital was the onset location for 233 (81%), a LTCF for 36 (13%), other healthcare facilities for 12 (4%) and unknown for five (2%).

Table 1. National CDI epidemiology: Q2 2018 versus Q2 2017.

| CDI case type | National CDI Epidemiology Q2 2018 vs Q2 2017            | Q2 2018 | Q2 2017 |
|---------------|---|---------|---------|
|               | Total reported cases:                                   | 527     | 451     |
|               | New   | 449     | 382     |
|               | Recurrent   | 48      | 41      |
|               | Unknown   | 30      | 28      |
| CDI origin    | Healthcare-associated (HCA):                            | 308     | 258     |
|               | Reporting hospital                                      | 250     | 207     |
|               | Long term care facility (LTCF)                          | 39      | 33      |
|               | Other healthcare facility                               | 17      | 18      |
|               | Unknown healthcare facility                             | 2       | -       |
|               | Community associated (CA)                               | 134     | 119     |
|               | Discharged within 4 – 12 weeks from healthcare facility | 46      | 31      |
|               | Unknown origin  | 39      | 43      |
| CDI onset     | Healthcare onset (HO):                                  | 286     | 237     |
|               | Reporting hospital                                      | 233     | 186     |
|               | LTCF  | 36      | 29      |
|               | Other healthcare facility                               | 12      | 13      |
|               | Unknown location  | 5       | 9       |
|               | Community onset (CO)                                    | 216     | 186     |
|               | Unknown onset location                                  | 25      | 28      |
| CDI severity  | Critical care admission or colectomy                    | 12      | 12      |

# Part 2: Hospital-acquired CDI (HA-CDI) Epidemiology – Q2 2018

Data on HA-CDI was reported from 55 acute public and private hospitals across Ireland. There were 250 HA-CDI cases in patients aged 2 years during Q2 2018. Of those, 221 were new HA-CDI cases, representing a national new HA-CDI rate of 2.2 [median rate = 1.4], higher than that reported for Q2 2017 [190 cases; rate = 1.9; median rate = 1.1]. *Figure 1* displays quarterly HA-CDI rates since 2010 and *Table 2* displays quarterly HA-CDI data from 2016 to 2018.



The national overall CDI rate represents all CDI diagnosed in hospitalised patients per 10,000 BDU, while the HA-CDI rate represents <u>new</u> cases of hospital-acquired CDI per 10,000 BDUs. Raw data for this graph is provided in Table 2. The national range is represented by the 5<sup>th</sup> to 95<sup>th</sup> percentile of the CDI rate.

Figure 1. Quarterly national HA-CDI rates: 2010 – 2018.

#### **CDI Case Type**

The majority of 250 hospital-acquired CDI cases were categorised as new infections (221; 88%), with 29 (12%) recurrent cases.

### **CDI Onset**

Patient locations at onset of HA-CDI symptoms included; while admitted to a healthcare facility, termed healthcare-onset (HO) for 218 cases (87%) and while residing in the community, termed community onset (CO) for 32 cases (13%).

Of 218 HO CDI cases, the reporting hospital was the onset location for 212 (97.2%), a long-term care facility for three cases (1.4%) and was unknown for three cases (1.4%).

Table 2. Quarterly HA-CDI data: 2016 – 2018

| YearQ  | Number of participating | Number of cases reported |           |         | CDI rate per 10,000 BDUs <sup>b</sup> |      |                    |        |
|--------|-------------------------|--------------------------|-----------|---------|---------------------------------------|------|--------------------|--------|
|        | hospitals <sup>a</sup>  | New                      | Recurrent | Unknown | Total                                 | Rate | Range <sup>c</sup> | Median |
| 2016Q3 | 51                      | 182                      | 22        | 2       | 206                                   | 2.0  | 0 - 3.8            | 1.4    |
| 2016Q4 | 52                      | 182                      | 19        | 2       | 203                                   | 2.0  | 0 - 3.3            | 1.2    |
| 2017Q1 | 51                      | 201                      | 20        | 2       | 223                                   | 2.0  | 0 - 3.8            | 1.4    |
| 2017Q2 | 54                      | 190                      | 16        | 1       | 207                                   | 1.9  | 0 - 3.5            | 1.1    |
| 2017Q3 | 55                      | 212                      | 27        | 0       | 239                                   | 2.2  | 0 - 4.2            | 1.5    |
| 2017Q4 | 56                      | 184                      | 26        | 4       | 214                                   | 1.8  | 0 - 3.8            | 1.2    |
| 2018Q1 | 55                      | 229                      | 18        | 0       | 247                                   | 2.2  | 0 - 5.1            | 1.3    |
| 2018Q2 | 55                      | 221                      | 29        | 0       | 250                                   | 2.2  | 0 - 3.9            | 1.4    |

a Since Q1 2012, 97% of all tertiary and general hospitals participated in the enhanced surveillance system.

# Part 3: C. difficile Testing Methods – Q2 2018

All 55 hospitals participating in the enhanced CDI surveillance system during Q2 2018 reported use of a *C. difficile* testing method recommended by the updated National Clinical Guidelines for Surveillance, Diagnosis & Management of *C. difficile* Infection in Ireland (2014). This includes either one of a variety of two-step testing methods (n=28; 51%) or a single-step method using molecular polymerase chain reaction (PCR) test for *C. difficile* toxin gene (n = 27; 49%) as displayed in *Table 3*, along with stratification by hospital type.

Table 3. C. difficile testing methods utilised in Q2 2018, by hospital type.

| Test Category  | Hospital Type |         |            |          | Total |
|--|---------------|---------|------------|----------|-------|
| rest category  | General       | Private | Specialist | Tertiary | Total |
| 1 STEP: PCR for toxin gene                                       | 12            | 3       | 6          | 6        | 27    |
| 2 STEP: GDH EIA, followed by confirmatory C. difficile toxin EIA | 3             | 4       | 0          | 0        | 7     |
| 2 STEP: Combined GDH with toxin EIA, followed by toxin EIA*      | 1             | 0       | 0          | 0        | 1     |
| 2 STEP: Combined GDH with toxin EIA, followed by PCR**           | 2             | 3       | 1          | 0        | 6     |
| 2 STEP: GDH EIA, followed by confirmatory PCR                    | 4             | 0       | 0          | 0        | 4     |
| 2 STEP: PCR, followed by confirmatory toxin EIA                  | 5             | 1       | 1          | 3        | 10    |
| Total  | 27            | 11      | 8          | 9        | 55    |

PCR for C. difficile toxin gene: Polymerase chain reaction (PCR) for the detection of TcdA and/or TcdB genes GDH EIA Enzyme immunoassay (EIA) for the detection of glutamate dehydrogenase (GDH) of C. difficile

**b** The CDI rate is the number of new cases of CDI that were acquired in the reporting hospital - per 10,000 bed days used (BDUs).

**c** The national range corresponds to the 5<sup>th</sup> to 95<sup>th</sup> percentile of the data. *Data for Q2 2018 is provisional* 

GDH AND TOXIN EIA: Enzyme immunoassay (EIA) for the detection of both *C. difficile* GDH and *C. difficile* toxin TcdA and/or TcdB

<sup>\*2</sup> STEP: Combined GDH with toxin EIA, followed by confirmatory toxin EIA: Addition of a confirmatory toxin EIA test (using a different EIA kit) if the initial toxin EIA is negative

<sup>\*\*2</sup> STEP: Combined GDH with toxin EIA, followed by confirmatory PCR: Addition of confirmatory PCR if the initial toxin EIA is negative

# Part 4: C. difficile Ribotyping - Q2 2018

Ribotyping data was available for just 20% of CDI cases reported to CDI enhanced surveillance, a reflection on the continued absence of a national funded *C. difficile* reference laboratory service, which has been a key recommendation of national *C. difficile* guidelines since 2008. Ribotypes 002, 078 and 014 were the most frequently reported. The lack of a robust, prospective system to capture *C. difficile* typing data limits understanding of the epidemiology of this important healthcare-associated infection.

## **Acknowledgments**

The HPSC would like to sincerely thank all who have contributed to this report: Microbiology Surveillance Scientists, Infection Prevention and Control Nurses, Microbiology Laboratory Scientists, Clinical Microbiologists, along with all the staff of the Departments of Public Health across Ireland.

# **Appendix A: National CDI Enhanced Surveillance Participating Hospitals**

|                                  | ial CDI Ennanced Surveillance  |            |
|----------------------------------|--|------------|
| Hospital Group                   | Hospital Name  | Category   |
|                                  | Adelaide & Meath & National Children's Hospital, Tallaght                    | ·          |
|                                  | Coombe Women and Infant's University Hospital                                | Specialist |
|                                  | Midland Regional Hospital Portlaoise   | General    |
| Dublin Midlands                  | Midland Regional Hospital Tullamore  | General    |
|                                  | Naas General Hospital  | General    |
|                                  | St James's Hospital  | Tertiary   |
|                                  | St Luke's Hospital, Dublin   | Specialist |
|                                  | Cappagh National Orthopaedic Hospital, Dublin                                | Specialist |
|                                  | Mater Misericordiae University Hospital                                      | Tertiary   |
|                                  | Midland Regional Hospital Mullingar  | General    |
|                                  | National Maternity Hospital, Holles Street                                   | Specialist |
|                                  | Our Lady's Hospital, Navan   | General    |
| Ireland East Hospital Group      | Royal Victoria Eye & Ear Hospital, Dublin                                    | Specialist |
|                                  | St Columcille's Hospital, Loughlinstown                                      | General    |
|                                  | St Luke's General Hospital, Kilkenny   | General    |
|                                  | St Michael's Hospital, Dun Laoghaire   | General    |
|                                  | St Vincent's University Hospital   | Tertiary   |
|                                  | Wexford General Hospital   | General    |
|                                  | Beaumont Hospital  | Tertiary   |
|                                  | Cavan General Hospital   | General    |
| RCSI Hospital Group              | Connolly Hospital, Blanchardstown  | General    |
| 22, 22, 22, 23,                  | Louth County Hospital, Dundalk   | General    |
|                                  | Our Lady of Lourdes Hospital, Drogheda                                       | General    |
|                                  | Letterkenny General Hospital   | General    |
|                                  | Mayo General Hospital, Castlebar   | General    |
|                                  | Portiuncula University Hospital, Ballinasloe                                 | General    |
| Saolta Hospital Group            | Roscommon University Hospital  | General    |
|                                  | Sligo General Hospital   | General    |
|                                  | University College Hospital Galway   | Tertiary   |
|                                  | Bantry General Hospital  | General    |
|                                  | Cork University Hospital Group   | Tertiary   |
|                                  | Kerry General Hospital, Tralee   | General    |
|                                  | •  |            |
| South /South Wast Haspital Group | Lourdes Orthopaedic Hospital, Kilcreene, Kilkenny<br>Mallow General Hospital | Specialist |
| South/South West Hospital Group  | •  | General    |
|                                  | Mercy University Hospital, Cork  | General    |
|                                  | South Infirmary - Victoria University Hospital, Cork                         | General    |
|                                  | South Tipperary General Hospital, Clonmel                                    | General    |
|                                  | Waterford Regional Hospital  | Tertiary   |
|                                  | Croom Hospital   | Specialist |
|                                  | Ennis Hospital   | General    |
| UL Hospital Group                | Nenagh Hospital  | General    |
|                                  | St John's Hospital   | General    |
|                                  | University Hospital, Limerick  | Tertiary   |
|                                  | University Maternity Hospital  | Specialist |
|                                  | Aut Even, Kilkenny   | Private    |
|                                  | Beacon Hospital, Dublin  | Private    |
|                                  | Blackrock Clinic   | Private    |
|                                  | Bon Secours, Cork  | Private    |
|                                  | Bon Secours, Galway  | Private    |
| Private Hospitals                | Bon Secours, Glasnevin   | Private    |
|                                  | Bon Secours, Tralee  | Private    |
|                                  | Galway Clinic  | Private    |
|                                  | Mater Private, Dublin  | Private    |
|                                  | Mater Private, Cork  | Private    |
|                                  | St Vincents Private Hospital   | Private    |
| Children's Hospital Group        | Children's University Hospital, Temple Street                                | Specialist |

## **Appendix B**

## Case Definitions for Surveillance of Clostridium difficile Infection

For surveillance purposes, a confirmed Clostridium difficile infection (CDI) case is a patient two years or older, to whom one or more of the following criteria applies:

- Diarrhoeal\* stools or toxic megacolon, with either a positive laboratory assay for *C. difficile* toxin A (TcdA) and/or toxin B (TcdB) in stools or a toxin-producing *C. difficile* organism detected in stool via culture or other means.
- Pseudomembraneous colitis (PMC) revealed by lower gastrointestinal endoscopy.
- Colonic histopathology characteristic of *C. difficile* infection (with or without diarrhoea) on a specimen obtained during endoscopy, colectomy or autopsy.
- \* Diarrhoea is defined as three or more loose/watery bowel movements (which are unusual or different for the patient) in a 24 hour period

#### CASE TYPE

- New Case of CDI:
  - The first episode of CDI, OR
  - A subsequent episode of CDI with onset of symptoms more than eight weeks after the onset of a previous episode.
- Recurrent Case of CDI:
  - A patient with an episode of CDI that occurs within eight weeks following the onset of a
    previous episode provided that CDI symptoms from the earlier episode resolved with or
    without therapy.

#### **ONSET**

- Healthcare onset » Symptoms start during a stay in a healthcare facility.
- Community onset » Symptoms start in a community setting, outside healthcare facilities.
- No information available » If no information was available on onset of symptoms

#### **ORIGIN**

- Healthcare-associated case. This is a CDI patient with either:
  - Onset of symptoms at least 48 hours following admission to a healthcare facility (healthcare-onset, healthcare-associated), OR
  - With onset of symptoms in the community within four weeks following discharge from a healthcare facility (community-onset, healthcare-associated).
- Community-associated case. This is a CDI patient with either:
  - Onset of symptoms while outside a healthcare facility, and without discharge from a healthcare facility within the previous 12 weeks (community-onset, community-associated), OR
  - With onset of symptoms within 48 hours following admission to a healthcare facility without residence in a healthcare facility within the previous 12 weeks (healthcare-onset, community-associated).
- Discharged 4 12 weeks from a healthcare facility
- »This is a CDI patient who was discharged from a healthcare facility between four and 12 weeks before the onset of symptoms.
  - No information available

#### **SEVERE CDI Case**

This is a CDI patient to whom any of the following criteria apply:

 Admission to an intensive care unit for treatment of CDI or its complications (e.g., for shock requiring vasopressor therapy)

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- Surgery (colectomy) for toxic megacolon, perforation or refractory colitis
- Death within 30 days after diagnosis if CDI is either the primary or a contributive cause