The first Canadian pediatric case of extensively drug-resistant \textit{Salmonella} Typhi originating from an outbreak in Pakistan and its implication for empiric antimicrobial choices

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\begin{abstract}
We report on a three-year-old male who contracted enteric fever during a visit to the Sindh province of Pakistan in the summer of 2018. He was diagnosed after returning to Canada and blood cultures isolated \textit{Salmonella enterica} serovar Typhi which harbored extensive drug-resistance (XDR) to all first-line antibiotics including ceftriaxone. Empiric ceftriaxone was switched to meropenem and he was successfully treated with a two-week course. An outbreak of XDR typhoid is currently emerging from Pakistan and several outbreak-related cases have been identified in the U.K and U.S. Whole genome sequencing confirmed that our child was infected with the XDR outbreak-strain. Current empiric antimicrobial choices will result in treatment failure if an XDR strain is encountered, therefore clinicians must adapt their empiric approach for those returning from high risk regions. This is the first XDR typhoid case in Canada and the first pediatric case to be diagnosed and treated outside of Pakistan. Clinicians must be vigilant for future cases.

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\section*{Introduction}

Enteric fever remains a significant disease that inflicts a great health burden worldwide, with an estimate of 21.6–26.9 million cases and 216 000 deaths each year attributed to \textit{Salmonella} Typhi \cite{1,2}. The current empiric management of typhoid in many centers is the use of third-generation cefalosporins. In November 2016, a large outbreak of extensively drug-resistant (XDR) typhoid has emerged in Pakistan which demonstrated resistance to all first-line antimicrobials including third-generation cefalosporins, thus creating a new challenge to empiric strategies.

\section*{Case report}

A healthy three-year-old boy became unwell during a holiday to Karachi, Pakistan in June-July 2018. Before his return to Canada, he developed fever, abdominal pain, diarrhea, and vomiting for two days. He attended a local physician, who prescribed cefixime and advised the family to follow up at the Emergency Department (ED) in Canada. There were no known infective contacts and he stayed with maternal grandparents in Karachi throughout the visit. The family only visited relatives and attended a family wedding in the same area. They consumed locally prepared food and drank bottled water. The family did not seek pre-travel advice and did not receive typhoid vaccine, malaria prophylaxis, or other travel related medication.

The child was seen at the ED at Toronto's Hospital for Sick Children soon after returning to Canada and was diagnosed with presumed enteric fever, empirically started on ceftriaxone and admitted to a nearby pediatric unit for further management. Stool and blood cultures obtained during the initial ED visit subsequently grew \textit{Salmonella enterica} serovar Typhi (S. Typhi).

Antimicrobial susceptibility testing of the isolate identified resistance to chloramphenicol, ampicillin, trimethoprim-sulfamethoxazole (TMP-SMX), ciprofloxacin and ceftriaxone but susceptible to meropenem and azithromycin (Table 1). After consultation with the Pediatric Infectious Disease service, the

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Table 1

| Antibiotic | MIC (mg/L) | Susceptibility |
|------------|------------|----------------|
| Chloramphenicol | >32 | Resistant |
| Ampicillin | >16 | Resistant |
| TMP-SMX | ≥320 | Resistant |
| Ciprofloxacin | 2 | Resistant |
| Ceftriaxone | >32 | Resistant |
| Meropenem | ≤0.125 | Sensitive |
| Azithromycin | ≤16 | Sensitive |
| Tigecycline | 0.25 | No interpretive guideline |

**Abbreviations:** MIC (minimal inhibitory concentration); TMP-SMX (Trimethoprim-sulfamethoxazole).

MIC interpretative criteria are based on Clinical and Laboratory Standards Institute M100-S28.

A novel patient was switched to intravenous meropenem at a dose of 20 mg/kg/dose on day two of admission. Blood cultures remained positive twenty-four hours after starting meropenem, but the cultures taken three days later were negative. Despite being treated with appropriate antibiotics, the child continued to have daily high fever for thirteen days into meropenem treatment albeit less frequent. Abdominal ultrasound scans only demonstrated mild hepatosplenomegaly which resolved and daily full-system examinations were unremarkable. He completed fourteen days of meropenem following which he was afebrile and clinically well. Three stool cultures obtained after completion of meropenem therapy were negative.

This patient’s S. Typhi isolate (S. Typhi strain PHLS950) underwent whole genome sequencing which was assessed using the ResFinder, SPIFindertools, SeqSero-1.2 Server, and PlasmidFinder [3]. Initially, the raw Illumina reads were trimmed generating 3,625,048 high quality reads corresponding to 886,576,883 detected bases and assembled using de novo assembler in CLC Genomics Workbench version 8.0.1 (CLC bio, Germantown, MD, USA) providing 85 contigs (accession no. RHPM000000000). The closest reference using functional comparison on the RAST server was found to be *Salmonella enterica subspecies enterica* serovar Typhi Ty2 reference genome NCBI: txisd2009261 (score, 534). BLAST result of the longest contigs (488765 bp) showed 100% identity to *S. enterica* subsp. *enterica* serovar Typhi (accession no. LT882486.1), a recently identified extensively drug-resistant S. Typhi strain from an outbreak in Pakistan, which encodes a chromosomally located resistance region and harbours a plasmid encoding additional resistance elements, including the *blaCTX-M-15* extended-spectrum β-lactamase, and carrying the *qnrS* fluoroquinolone resistance gene (Fig. 1). The ResFinder tool identified multiple resistance genes conferring resistance to various antibiotics (Supplementary Table 1). In addition, mutation at 583 of the *gyrA* gene was also identified. Based on the PlasmidFinder tool, we identified two plasmids: IncY plasmid, which matched 100% to the plasmid that was isolated in the Pakistan outbreak-strain (accession no. LT906492) [4], and IncQ1 plasmid.

**Discussion**

To our knowledge, this is one of the first cases of XDR typhoid in a child outside of Pakistan. Typhoid fever caused by *S. Typhi*, continues to inflict a significant health burden worldwide with reports estimating 216–26.9 million cases and 216 000 typhoid-associated deaths each year [1,2]. The highest incidence of typhoid is in low to middle income countries (LMIC) that have poor sanitation and public health infrastructure; rates are highest in South Asia [5].

The emergence of *S. Typhi* strains resistant to chloramphenicol first appeared in the early 1970s [6] and resistance to other first line agents including ampicillin and TMP-SMX became increasingly prevalent in the 1980s and early 1990s, leading to the term multidrug resistant (MDR) typhoid. Fluoroquinolones, particularly ciprofloxacin, become first line therapy in the 1990s however, resistance developed shortly thereafter. Phylogenetic studies identified MDR typhoid to be associated with the dominant H58 lineage which has spread throughout the world, namely Asia and Africa [7]. The mode of resistance acquisition is through a combination of plasmid transfer and integrated antimicrobial resistance genes at different chromosomal loci [4,6,7]. Due to the emergence of MDR typhoid, most nations are now reliant upon third-generation cephalosporins like ceftriaxone and azidilites like azithromycin, as first-line agents. For LMIC, these antimicrobials have become the only financially feasible options.

International travel is increasing, thus creating a need for clinicians to be vigilant of diseases that are commonly imported from abroad and of the resistance profiles that these infections may harbor. Travelers who visit friends and relatives (VFRs), particularly to the Indian subcontinent, are at the highest risk of contracting enteric fever [8]. In a recent twenty-eight-year review of all children with enteric fever who presented to Toronto’s Hospital for Sick Children, 83% were VFRs and 80% were acquired during travel to Pakistan, India or Bangladesh [8]. All isolates in this study were sensitive to ceftriaxone, which is consistent with the common practice in Ontario to treat suspected typhoid empirically with ceftriaxone.

Over the last two decades, sporadic cases have been reported of XDR typhoid resistant to ceftriaxone [4] and even azithromycin [9]. Since November 2016, Pakistan’s Sindh province has experienced a large outbreak of XDR *S. Typhi*, predominantly in the Hyderabad and Karachi areas. As of December 2018, more that 5000 cases have been reported and all isolates have been resistant to ampicillin, TMP-SMX, ciprofloxacin and ceftriaxone but remain susceptible to azithromycin [4,10,11]. The Sindh-outbreak strain is associated with the H58 haplotype and holds a chromosomally integrated composite transposon at the *yidA* locus which carries resistant genes to chloramphenicol (*catA1*), ampicillin (*blaTEM-1*), TMP-SMX (*dfrA7, sul1, sul2*) and ciprofloxacin (*chromosomal gyrA mutation* 583 F). This strain also includes an IncY plasmid that harbors a resistance gene to ciprofloxacin (*qnrS*) and an extended-spectrum β-lactamase (ESBL) gene conferring resistance to ceftriaxone (*blaCTX-M-15*). The MDR H58 haplotype likely transformed to the XDR strain after acquiring the ESBL harboring IncY plasmid from an *Escherichia coli* in Pakistan [4]. Our strain as well as the isolated plasmid are identical to the Sindh-outbreak strain.

So far in 2018, there have been six Pakistan outbreak-related cases of XDR typhoid in travelers from the United Kingdom and United States [4,6,11]. With the ease of travel, this number will certainly increase and will have implications for the clinicians who look after returning travelers from outbreak areas, namely choice of empiric antibiotics. The current practice of initiating ceftriaxone in these patients will be ineffective if the infecting strain is XDR. A potential empiric strategy is the use of meropenem for the septic child or the combination of ceftriaxone and azithromycin for the clinically stable, with further rationalization once sensitivities become available.

A recent significant advancement in the containment of typhoid is the prequalification of the Vi polysaccharide tetanus–toxoid conjugate vaccine (Tybar–TCV™) issued by the World Health Organization (WHO) in March 2018. The WHO recommends implementation of large scale TCV vaccination programs in endemic areas, which is effective from six months of age [12,13]. We report the first XDR typhoid case in Canada diagnosed in a 3 year old child returning from Pakistan, who was treated successfully with two weeks of meropenem. Clinicians must be vigilant of typhoid resistance in high-risk areas such as Pakistan and develop new strategies for empiric management.
Conflict of interest

No conflict of interest to declare by authors.

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Ethical approval

Ethical approval was not required for this work.

Consent

Written informed consent was obtained from the patient for publication of this case report. A copy of the written consent is available for review by the Editor-in-Chief of this journal on request.

Author contribution

Waison Wong: First author with main role in writing the original draft.
Hatem Al Rawahi: Assisted in writing the original draft.
Samir Patel: Assisted in writing and carried out whole genome sequencing.
Yvonne Yau: Assisted in writing (review and editing).
Alireza Eshaghi: Assisted in writing and carried out whole genome sequencing.
Sandra Zittermann: Assisted in writing and carried out whole genome sequencing.
Leah Tatum: Assisted in writing (review and editing).
Shaun Morris: Supervision, writing (review and editing).

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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.idcr.2019.e00492.

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Fig. 1. Graphical circular map of S. Typhi PHL 5950 de-novo assembly contigs compare to strains LT882486.1 and strain Ty2 drawn with CGView. S. Typhi strain Ty2 used as reference for coding regions.
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