A Case Series of Diphtheria in Western India: Time to Revisit Vaccination Coverage and Policy?

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Abstract

INTRODUCTION: Diphtheria is a significant child health problem in countries with low immunization coverage. Reports of diphtheria in the adult population are also increasing. Here we describe case series of diphtheria in western India for 6 months. OBJECTIVE: To identify and isolate Corynebacterium species, demonstrate the isolates for toxigenicity and examine the immune status of the patients by estimating antibody titers in sera (anti-diphtheria toxoid IgG). METHODS: Twelve patients admitted with clinical suspicion of oropharyngeal diphtheria for six months. In each case, two throat swabs were collected and primary identification of Corynebacterium diphtheriae was done by direct microscopy (Gram's and Albert's stains), bacteriological culture and biochemical tests as per the standard procedure. Culture isolates were tested for toxigenicity by Elek's gel precipitation and were sent to a reference laboratory for tox A gene detection by polymerase chain reaction. Anti-diphtheria toxoid IgG antibody levels were determined in patient’s sera using a commercial Anti-Diphtheria Toxoid IgG Enzyme-Linked Immunosorbent Assay (EUROIMMUN, Germany) at a reference laboratory. RESULTS: All 12 patients presented with oropharyngeal diphtheria with the formation of pseudomembrane in the oropharynx. Eleven patients were of the pediatric age group and one was an adult. The microbiological diagnosis was achieved for 11 patients and one was diagnosed clinically. Based on vaccination history, microbiological findings and distribution of anti-diphtheria toxoid IgG antibodies titers, the results showed persistence of toxigenic strain of Corynebacterium diphtheriae circulating in our region. CONCLUSIONS: The present study demonstrated that toxigenic strains of C. diphtheriae are circulating in this geographical location which indicates the need for constant epidemiological surveillance ensuring early detection of diphtheria and review the efficacy of the immunization programme.

Keywords: Corynebacterium diphtheriae, tox A gene, immunization

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1. Introduction

Diphtheria, caused by toxin-producing strains of Corynebacterium diphtheriae, uncommonly Corynebacterium ulcerans and Corynebacterium pseudotuberculosis, is a severe disease with a fatality of 5 to 10%. In developing countries like India, diphtheria remains a significant child health problem with an increase in notified cases of diphtheria in the adult population as well [1].

Humoral immunity protects by forming neutralizing IgG antibodies to diphtheria toxin, which may be induced by either natural infection, passive or active immunization [2]. Following the success of the Universal Immunization Programme, diphtheria cases in India fell from nearly 40,000 in 1980 to 2,300 in 2015 [3]. However, due to reduced opportunities for booster immunization through subclinical infections, the levels of antibodies decreased with time leading to re-emergence of cases, also making adults susceptible to diphtheria and impacting the overall immunity of the population [4].

To illustrate the phenomenon of waning immunity to diphtheria in our population, we present a case series of diphtheria from a tertiary care center in Western India along with vaccination history and antibody titers (anti-diphtheria toxoid IgG) of the patients.

2. Objective

To identify and isolate Corynebacterium species, demonstrate the isolates for toxigenicity and examine the immune status of the patients by estimating antibody titers in sera (anti-diphtheria toxoid IgG).

3. Methods

This case series examined 12 patients admitted to our tertiary care center in South Maharashtra, India with clinical suspicion of oropharyngeal diphtheria for six months. This study was approved by the Institutional Ethics Committee (Dr. Vaishampayan Government
Medical College, Solapur. Ref no: Pharma Dept/IEC/Approval letter/121/21) and an informed written consent was obtained from all patients enrolled in this study. Care has been taken to ensure that participant’s identifiers are removed and identities are not revealed. In each case, two throat swabs were collected and primary identification of *Corynebacterium diphtheriae* was done by direct microscopy (Gram’s and Albert’s stains, Figure 2 and Figure 3), bacteriological culture, and biochemical tests as per the standard procedure [5]. Culture isolates were tested for toxigenicity by Elek’s gel precipitation test and were sent to a reference laboratory for *tox A* gene detection by polymerase chain reaction.

Patient sera were collected (before administration of anti-diphtheritic serum) and anti-diphtheria toxoid IgG antibody titers were determined at a reference laboratory using a commercial Anti-Diphtheria Toxoid IgG Enzyme-Linked Immunosorbent Assay (EUROIMMUN, Germany) [6]. Titters were interpreted as follows:

- <0.01 IU/ml = No protection.
- 0.01 – 0.099 IU/ml = Uncertain protection.
- ≥0.1 IU/ml = Immunization protection present.
- 1.0 IU/ml = Long-term immunization protection.

**4. Results**

All 12 patients presented with oropharyngeal diphtheria with the formation of pseudomembrane in the oropharynx. Eleven patients were of the pediatric age group and one was an adult. Of the 12 patients, eight (66.66%) recovered, three (25%) succumbed and one (8.33%) was lost to follow up (discharged against medical advice). Their vaccination histories, microbiological findings, distribution of anti-diphtheria toxoid IgG antibodies and outcomes are presented in Table 1. The microbiological diagnosis was achieved for 11 patients and one was diagnosed clinically. Six patients (50%) had levels of anti-diphtheria toxoid IgG antibodies below 0.01 IU/ml. Two patients (17%) showed uncertain protection (0.01 - 0.099 IU/ml), and the remaining four patients were not tested for antibody titers as they were administered with diphtheria antitoxin immediately on admission before blood samples could be collected due to their severe presentation. There was no significant difference in diphtheria antibody levels between males and females.

**5. Discussion**

Diphtheria is a fatal but vaccine-preventable disease, the incidence of which had declined worldwide following global immunization programme. With the inclusion of the diphtheria vaccine in the national immunization schedule, India saw a decline in the number of annual cases from 100,000 in 1980 to 2,500 in 2015. However, India accounted for a substantial proportion of the global burden of diphtheria cases [1]. In recent years, a resurgence was noted primarily due to low vaccine coverage and waning adult immunity to the disease. In our series, all cases except one were of the pediatric age group; either unimmunized (no DPT dose) or partially immunized (1 dose of DPT). None of the patients of the pediatric age group completed primary immunization. One adult case notified with complete primary immunization history but no booster immunization. Nath B et al [7] and Saikia L et al [8] also showed maximum cases in the adult
The World Health Organization recommends 3 primary doses (by six months of age) followed by 3 booster doses (by adolescence) of the diphtheria vaccine before adolescence. Diphtheria is a resurgent problem and a significant age shift towards adults has been notified in some regions [9,10]. However, the information about booster doses is limited and is expected to be low as the data is not routinely collected through surveys [1].

As demonstrated by the last major diphtheria epidemic in the former Soviet Union, when more than 50,000 cases were recorded, waning immunity due to vaccination and reduced natural infection leads to a large pool of susceptible persons, creating an epidemic potential. During this epidemic adolescents and adults were mainly affected, with a history of previous vaccination in most individuals [11]. Over the last 20 years (i.e. 1996-2016), hospital-based sentinel surveillance and outbreaks that were published in the past showed that diphtheria cases were most commonly reported in school going children and adolescents [1].

A survey from India found that 29% of 8,309 children aged 5-17 years were immune, 59.8% partially immune, and 10.5% non-immune, with variations between genders, regions and urban and rural settings. The same study also reported vaccine coverage (of the three primary doses) of 78.4% [12].

India reported a diphtheria-tetanus and pertussis (DTP3) vaccine coverage of nearly 90% among 1-year olds in 2019 versus a global coverage of 85%. Studies conducted worldwide and in India have documented the determinants of vaccination coverage and have catalogued the strategies that have proven effective in improving immunization coverage [13,14,15]. However, the diphtheria vaccine prevents disease but not carriage or transmission [16]. Our study showed, six patients (50%) had levels of anti-diphtheria toxoid IgG antibody titers below 0.01 IU/ml and two patients (17%) showed uncertain protection (0.01-0.099 IU/ml), thus measuring antibody titers against diphtheria toxin in individuals is the only way to survey the level of protection in a community and would also help to know immunization status. The limitation of the present study was four patients out of 12 were not tested for serum antibody titers as they were given Anti-Diphtheria serum (ADS) before blood sample collection.

### Table 1. Distribution of vaccination history, microbiological findings, distribution of anti-diphtheria toxoid IgG antibody titers and patient outcome

| CASE (AGE/SEX) | VACCINATION HISTORY | MICROSCOPY AND CULTURE | SERUM ANTI-DIPHTHERIA TOXOID IgG (IU/ML) | TOX A GENE | ANTI-DIPHTHERIA SERUM | OUTCOME | COMPLICATION |
|----------------|----------------------|-------------------------|-----------------------------------------|------------|----------------------|---------|--------------|
| PATIENT 1 11YRS/F | PARTIALLY VACCINATED* | POSITIVE | NOT AVAILABLE | POSITIVE | GIVEN | RECOVERED | - |
| PATIENT 2 7YRS/F | UNVACCINATED | POSITIVE | NOT AVAILABLE | POSITIVE | GIVEN | DEATH | MYOCARDITIS |
| PATIENT 3 10YRS/M | PARTIALLY VACCINATED* | POSITIVE | NOT AVAILABLE | POSITIVE | GIVEN | RECOVERED | - |
| PATIENT 4 7YRS/M | PARTIALLY VACCINATED* | POSITIVE | NOT AVAILABLE | POSITIVE | GIVEN | RECOVERED | - |
| PATIENT 5 8YRS/M | UNVACCINATED | POSITIVE | <0.01 | POSITIVE | GIVEN | RECOVERED | - |
| PATIENT 6 6YRS/M | PARTIALLY VACCINATED* | POSITIVE | <0.01 | POSITIVE | GIVEN | DEATH | MYOCARDITIS |
| PATIENT 7 7YRS/M | PARTIALLY VACCINATED* | POSITIVE | 0.01 - 0.099 | POSITIVE | GIVEN | RECOVERED | - |
| PATIENT 8 21YRS/M | VACCINATED (NO BOOSTER DOSE) | POSITIVE | <0.01 | POSITIVE | GIVEN | RECOVERED | - |
| PATIENT 9 10YRS/F | PARTIALLY VACCINATED* | POSITIVE | <0.01 | POSITIVE | NOT GIVEN | LOST TO FOLLOW UP | - |
| PATIENT 10 8YRS/M | PARTIALLY VACCINATED* | NEGATIVE (CLINICAL DIAGNOSIS) | 0.01 - 0.099 | NA | NOT GIVEN | RECOVERED | - |
| PATIENT 11 6YRS/F | UNVACCINATED | POSITIVE | <0.01 | POSITIVE | GIVEN | DEATH | - |
| PATIENT 12 6YRS/F | UNVACCINATED | POSITIVE | <0.01 | POSITIVE | GIVEN | RECOVERED | - |

*Only single dose received.
F = Female, M = Male, yrs = years in age.
Our series highlights the need to:

1. Improve primary vaccine coverage.
2. The immunity of school-going children could be improved by administering all the children with adult type of tetanus and diphtheria (Td) vaccine at school entry and replacing the tetanus toxoid vaccine used in school health program with Td vaccine; since the full potency diphtheria toxoid (25 Lf per dose) used in the Universal Immunization Program is associated with high reaction rates among older children and adults [1].
3. The use of Td instead of TT is to be recommended during pregnancy not only to protect against maternal and neonatal tetanus but also protect against diphtheria during prenatal care. Though implemented in some places, there is a need to spread awareness among people with low educational status.
4. Good surveillance is needed to document the impact of vaccination.
5. National level surveys need to include information about diphtheria boosters, besides the information about primary vaccination.
6. Conduct serosurveys in all the states covering different age groups to estimate the population immunity guide the immunization program.

6. Conclusion

The present study demonstrated that toxigenic strains of *C. diphtheriae* are circulating in this geographical location which indicates the need for constant epidemiological surveillance ensuring early detection of diphtheria and review the efficacy of the immunization programme. Detection of Diphtheria cases in adults – is an eye-opener for Physicians and epidemiologists, studies should be carried out with larger sample size. Due to the low levels of anti-diphtheria antibodies in adolescent and increased travel to diphtheria endemic regions, it seems reasonable to carry out a publicity campaign regarding recommendations for diphtheria booster vaccination in adults. Full protection in the highest possible proportion of the population should help to avoid the re-emergence of this serious, potentially fatal infectious disease.

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