RESEARCH ARTICLE

Antibiotics resistance in El Tor *Vibrio cholerae* 01 isolated during cholera outbreaks in Mozambique from 2012 to 2015

Liliana Candida Dengo-Baloi*, Cynthia Amino Sema´-Baltazar*, Lena Vania Manhique*, Jucunu Elias Chitio*, Dorteia Luı´sa Inguane*, Jose´ Paulo Langa*

Instituto Nacional de Saúde, Maputo, Moçambique

* These authors contributed equally to this work.

liliana.deng o.baloi@gmail.com, liliana.baloi@ins.gov.mz

Abstract

Rationale

Mozambique has recorded cyclically epidemic outbreaks of cholera. Antibiotic therapy is recommended in specific situations for management and control of cholera outbreaks. However, an increase in resistance rates to antibiotics by *Vibrio cholerae* has been reported in several epidemic outbreaks worldwide. On the other hand, there are few recent records of continuous surveillance of antibiotics susceptibility pattern of *V. cholerae* in Mozambique.

Goals

The purpose of this study was to evaluate antibiotics resistance pattern of *Vibrio cholerae* O1 Ogawa isolated during Cholera outbreaks in Mozambique to commonly used antibiotics.

Methodology

We analyzed data from samples received in the context of surveillance and response to Cholera outbreaks in the National Reference Laboratory of Microbiology from the National Institute of Health of Mozambique, 159 samples suspected of cholera from cholera treatment centers of, Metangula (09), Memba (01), Tete City (08), Moatize (01), Morrumbala (01) districts, City of Quelimane (01), Lichinga (06) and Nampula (86) districts, from 2012 to 2015. Laboratory culture and standard biochemical tests were employed to isolate and identify *Vibrio cholerae*; serotypes were determined by antisera agglutination reaction in blade. Biotype and presence of important virulence factors analysis was done by PCR. Antibiotics susceptibility pattern was detected by disk diffusion method Kirby Bauer. Antibiotic susceptibility and results were interpreted by following as per recommendations of CLSI (Clinical and Laboratory Standards Institute) 2014. All samples were collected and tested in the context of Africhol Project, approved by the National Bioethics Committee for Health.
Results

Among isolates from of *Vibrio cholerae* O1 El Tor Ogawa resistance to Sulphamethoxazole-trimethropim was 100% (53/53) to Trimethoprim-, being 100% (54/54) for Ampicillin, 99% (72/74) for Nalidixic Acid, 95% (42/44) for Nitrofurantoin and (19/20) Cotrimoxazole, 83% (80/97) Tetracycline, 56% (5/13) Doxycycline, 56% (39/70) Azithromycin and 0% (0/101) for Ciprofloxacin. PCR analysis suggested strains of *V. cholerae* O1 being descendants of the current seventh pandemic *V. cholerae* O1 CIRS 101 hybrid variant. The *V. cholerae* O1 currently causing cholera epidemics in north and central Mozambique confirmed a CTXΦ genotype and a molecular arrangement similar to the *V. cholerae* O1 CIRS 101.

Conclusion

Although *V. cholerae* infections in Mozambique are generally not treated with antibiotics circulating strains of the bacteria showed high frequency of in vitro resistance to available antibiotics. Continuous monitoring of antibiotic resistance pattern of epidemic strains is therefore crucial since the appearance of antibiotic resistance can influence cholera control strategies.

Introduction

*Vibrio cholerae* toxin is the virulence factor causing cholera disease, which is characterized by a secretory acute diarrhea. Cholera can lead to severe dehydration and death within hours if not promptly treated. Cholera constitutes a serious public health problem in many parts the world [1,2].

The seventh pandemic of cholera reached Africa in the 70’s, arrived in Mozambique in 1973 with cases reported until 1985. Cholera resurfaced in 1989 with over 3,600 cases reported, caused by *V. cholerae* O1 El Tor serotype Inaba (with some cases of Ogawa), susceptible to Tetracycline, Chloramphenicol and sulfadiazine. [3].

Mozambique continued on experiencing recurrent outbreaks of cholera, in different parts of the country, with different spatial pattern from year to year. Increase in antibiotic resistance including resistance to recommended antibiotic for treatment has also been reported [4,5]. The last decades have seen a growing trend in antimicrobial resistance in Mozambique, previous researches have reported *V. cholerae* Ogawa O1 El Tor drug resistance in the last decade for Maputo [6–6], Zambezia and Tete [5].

Antimicrobial resistance of *V. cholerae* O1 El Tor is of interest because it’s becoming a serious public health problem for many African countries [7]. Therapy with effective antimicrobial agents significantly reduces the duration of diarrhea and hospitalization, reduces the volume of watery feces and need for maintenance fluids. The duration of fecal excretion of *V. cholerae* is also decreased, reducing transmission of infection to family members, as well as nosocomial infections [8].

Increasing drug resistance is well known and usually varies from one place to another. *V. cholerae* becomes drug resistant by exporting drugs through efflux pumps, chromosomal mutations or developing genetic resistance via the exchange of conjugative plasmids, conjugative transposons, integrons or self-transmissible chromosomally integrating *sxt* elements. In addition, *V. cholerae*, as an environmental organism, have means to acquire resistance genes.
from intimate contact with intrinsically resistant environmental bacteria, through mobile genetic elements. *V. cholerae* is able to share these antibiotic resistance genes with other bacteria; and once in the human gut, the bacteria may share these resistance traits with commensals or other enteric pathogens, what complicates antibiotic therapy of many infections [2].

Since antibiotic therapy is recommended in specific situations for management and control of cholera outbreaks, monitoring *V. cholerae* resistance is important for public health. The aim of this study was to describe spatiotemporally the antimicrobial resistance pattern of *Vibrio cholerae* O1 El Tor Ogawa isolated in patients admitted to cholera treatment centers (CTCs) and diarrheic disease treatment centers (CTDDs) during outbreak investigations in Mozambique, from 2012 to 2015.

**Material and methods**

**Sample collection and transportation**

Rectal swabs from suspected cholera cases were collected during Cholera outbreaks from CTCs and CTDD’s from 2012 to 2015 by local laboratory technicians and transported to local laboratories in Cary Blair medium, prepared from dehydrated media. These samples were collected after patient’s stabilization and before antimicrobial administration.

**Culture and identification**

Laboratory technicians, at provincial level, when possible, *Vibrio cholerae* were identified by standard culture methods; and sent to National Reference Laboratory of Microbiology (NRLM), at central level, for serological confirmation, Antimicrobial susceptibility test (AST) and a double multiplex PCR for mobilome profile analysis.

In the laboratory, samples were enriched in APA broth, then cultured in TCBS media, and suspected colonies were submitted to standard biochemical tests; when positive, colonies were submitted to serological tests for *Vibrio cholerae* TM Difco BD Poly and *Vibrio cholerae* TM Difco BD Ogawa (Denka Seiken, Tokyo, Japan).

**Antimicrobial susceptibility testing (Ast)**

Serologically positive samples were submitted to AST by Kirby Bauer diffusion method [9] for commonly used and recommended antibiotics, namely, Nalidixic acid, Ampicillin, Sulphamethoxazole-trimethoprim (SXT), Tetracycline, Chloramphenicol, Nitrofurantoin, Azithromycin, Cotrimoxazole and Doxycline, according to laboratory’s antibiotic discs availability at the moment. Results from AST were interpreted using CLSI (2014).

**PCR analyses**

Representatively, positive isolates on serological tests were submitted to a double multiplex PCR essay, using five completely sequenced reference strains as positive controls (MJ1236, N1696, B33, CIRS101 and MO10), for the presence of RstR, ctxB, and tcpA genes, genetic markers for El Tor Biotype [10].

**Ethical considerations**

Analyzed data was from samples collected during Africhol Project, the African Cholera Surveillance Network multi-centric project consisting of an 11 African countries consortium and non-governmental organizations, aiming to collect epidemiological and microbiological information for the occurrence of cholera in Africa to advise for control and preventive measures ([http://africhol.org/](http://africhol.org/)), which surveillance protocol was approved by the Mozambican National
Bioethics Committee for Health. During the study, all participants gave an informed written consent.

We analyzed Africhol data from the laboratory records, anonymously (all samples were codified upon entrance), there was no direct intervention or interaction with human subjects and neither identifiable private information.

Results

Mozambique reported 27 outbreaks in 22 districts (of 145), on 07 of 10 provinces (Fig 1), from 2012 to 2015; and 1522 samples of suspected Cholera cases were received at the National Reference Laboratory of Microbiology, 510 confirmed as *V. cholerae* serogroup O1 biotype El Tor serotype Ogawa. From these 510, 159 were submitted to AST.

Table 1 shows *V. cholerae* O1 El Tor Ogawa antibiotic resistance during epidemics, between 2012 and 2015. In 04 isolates tested on 2012’s outbreaks we found no resistance to Ciprofloxacin and Azithromycin, low levels of resistance to Tetracycline (25%), and 100% resistance to Ampicillin, Nalidixic Acid, Chloramphenicol, SXT and Doxycyclin.

On 2013, we tested 19 isolates with no resistance to Ciprofloxacin, low resistance to Doxycyclin (11%), increased resistance to Tetracycline (32%), resistance to Chloramphenicol (58%) and 100% resistance to Ampicillin, Nalidixic Acid, SXT and Azithromycin.

In 2014, for 35 isolates, we found no resistance to Ciprofloxacin and SXT, increased resistance to tetracycline (44%), high resistance to Nitrofurantoin (94%), Chloramphenicol (97%) and 100% to Ampicillin and Nalidixic acid.

For 2015 isolates (n = 100) we found 100% resistance for Ampicillin, Tetracycline, Nalidixic acid, Chloramphenicol, SXT and Nitrofurantoin; 95% resistance to Cotrimoxazole and no resistance to Ciprofloxacin.

Tendency of resistance from 2012 to 2015 (Fig 2) indicates an increase in antimicrobial resistance for Tetracycline, Nitrofurantoin, and Azithromycin. A sudden decrease and subsequent increase of resistance in Chloramphenicol and SXT, decrease in Doxycycline from 2012 to 2013, 95% resistance in 2015 for Cotrimoxazole and 100% resistance along the years for Ampicillin and Nalidixic Acid. Laboratory analyses also shows that *V. cholerae* O1 El Tor Ogawa isolated during cholera outbreaks in Mozambique, have no resistance to Ciprofloxacin.

Overall *V. cholerae* from 2012 to 2015 outbreaks had 100% resistance to Ampicillin and Nalidixic Acid, 97% to Nitrofurantoin, 95% to Cotrimoxazole, 89% to Chloramphenicol, 75% to SXT, 56% to Doxycycline, 50% to Tetracycline, 13% to Azithromycin and 0% to Ciprofloxacin.

Genetic screening by PCR in 58 isolates revealed three important El Tor epidemic markers, *ctxA*, *rstR2*, and *tcpA*, and the presence of CTX ϕ on chromosome 1 instead of chromosome 2, confirming the profile found in *V. cholerae* O1 El Tor variants B33 and CIRS 101. (Table 2)

Discussion

The current study found increasing antibiotic resistance in *Vibrio cholerae* O1 El Tor Ogawa isolated from Cholera outbreaks from 2012 and 2015 in Mozambique to Tetracycline, Trimethoprim-sulphamethoxazol, Chloramphenicol and Nitrofurantoin.

Increasing resistance to tetracycline was consistent with data from Zambia [11], urban and rural Bangladesh [12]; DRC [13]; Nigeria [14] and South Mozambique [4,6]. However, decreasing resistance to tetracycline have been reported in other places, such as Calcutta [15], East Delhi [16] and Ghana [17] and susceptibility to tetracycline, in Puduchuary in India [18], Haiti [19] and north India [20].
Fig 1. AST per district and per year, for *V. cholerae* O1 El Tor Ogawa isolated during cholera outbreaks in Mozambique from 2012 to 2015. In 2012 (Cuamba district in Niassa province and Montepuez district in Cabo Delgado province), 2013 (Cuamba district in Niassa province, Pemba city and Montepuez districts in Cabo Delgado province, Nampula city *district* in Nampula province and Alto-Molóque *district* in Zambezia province), 2014 (Nampula city *district* in Nampula province), 2015 (Lichinga city, Lago and Cuamba districts in Niassa province, Nampula city *district* in Nampula province and Morrumbala and Quelimane city *districts* in Zambèzia province, Tete city and Moatize districts in Tete province, Beira city *district* in Sofala province and Matola city *district* in Maputo province). AMP- Ampicillin; TE- Tetracycline; NA- Nalidixic Acid; C- Chloramphenicol; CIP- Ciprofloxacin; SXT- Sulphamethoxazole-trimethoprim; F- Nitrofurantoin; AZM- Azithromycin; rstR, ctxB and tcpA- *Vibrio cholerae* virulence genes; TLC-RS1, CORE-RTX and TCL-RS2—primers for the presence of CTXϕ on chromosome 1; Chr II—Chromosome 2

https://doi.org/10.1371/journal.pone.0181496.g001
Our results indicating resistance to Nitrofurantoin and SXT, match to those observed in earlier studies in Zambia [11], and DRC [13]; and SXT resistance also in a Mozambique rural area [4], and in north India [20].

SXT resistance pattern was in accord with Folgosa et al ten years ago in South and Central Mozambique, having along studied years, all isolates resistant to this antibiotic disc except for one, that was sensitive to it [5].

Another important finding was that isolates were 100% sensitive to Ciprofloxacin activity throughout the studied years; In agreement with Folgosa [5] in 3 provinces in Mozambique (Maputo, Zambezia and Tete), Gujral in Maputo city and province; although these results differ from some published studies in Calcutta [15], East Delhi [16], Nigeria [14], Haiti [19], Iran [21], urban and rural Bangladesh [12], Ghana [17] and Indonesia [22] with resistance or increasing resistance to Ciprofloxacin.

Along the years, isolates in our study were 100% resistant Ampicillin, which is consistent with results from Calcutta [15], Zambia [11] and in contrast with Mozambique in the last decade, North India [20] and Indonesia [22].

| Antimicrobial agent          | 2012   | 2013   | 2014   | 2015   | % (R) |
|-----------------------------|--------|--------|--------|--------|-------|
| Ampicillin (AMP)            | 10μg   | 100%   | 100%   | 100%   | 100%  |
| Tetracycline (TE)           | 30μg   | 25%    | 32%    | 44%    | 50%   |
| Nalidixic Acid (NA)         | 30μg   | 100%   | 100%   | 100%   | 100%  |
| Chloramphenicol (C)         | 10μg   | 100%   | 58%    | 97%    | 100%  |
| Ciprofloxacin (CIP)         | 5μg    | 0%     | 0%     | 0%     | 0%    |
| Trimethoprim-Sulphamethoxazole (SXT) | 23.75/1.25μg | 100% | 100% | 0% | 100% | 75% |
| Nitrofurantoin (F)          | 300μg  | n.t.   | n.t.   | 94%    | 100%  | 97%   |
| Azithromycin (AZM)          | 15μg   | 0%     | 0%     | n.t.   | 39%   | 13%   |
| Cotrimoxazol (CTX)          | 30μg   | n.t.   | n.t.   | n.t.   | 95%   | 95%   |
| Doxycyclin (DO)             | 30μg   | 100%   | 11%    | n.t.   | 56%   |

n.t. = not tested

https://doi.org/10.1371/journal.pone.0181496.t001

Fig 2. Percentage of antibiotic resistance in *V. cholerae* O1 El Tor Ogawa isolated during cholera outbreaks in Mozambique from 2012 to 2015. AMP- Ampicillin; TE- Tetracycline; NA- Nalidixic Acid; C- Chloramphenicol; CIP- Ciprofloxacin; SXT- Sulphamethoxazol-trimethoprim; F- Nitrofurantoin; AZM- Azithromycin.

https://doi.org/10.1371/journal.pone.0181496.g002
Table 2. CTX $\phi$ cluster analysis of *Vibrio cholerae* O1 El Tor Ogawa isolated during cholera outbreaks in Mozambique from 2012 to 2015. Showing a classic signature of *Vibrio cholerae* O1 El Tor variants B33 and CIRS 101. AMP- Ampicillin; TE-Tetracycline; NA- Nalidixic Acid; C-Chloramphenicol; CIP-Ciprofloxacin; SXT- Sulphamethoxazol-trimethoprim; F- Nitrofurantoin; AZM- Azithromycin; rstR, ctxB and tcpA- *Vibrio cholerae* virulence genes; TLC-RS1, CORE-RTX and TCL-RS2—primers for the presence of CTX $\phi$ on chromosome 1; Chr II—Chromosome 2.

| Year | Tested isolates | Origin | AMP | TE | NA | C | CIP | SXT | 23.75/1.25ug | ICE | VSP-II | TLC | Kappa | GI-12 | GI-14 | GI-15 | Profile |
|------|-----------------|--------|-----|----|----|---|-----|-----|-------------|-----|--------|-----|-------|-------|-------|-------|---------|
| 1975 | N16961          | India  | -   | -  | -  | - | -   | -   | -           | +   | -      | -   | -     | -     | -     | -     | A       |
| 1992 | MO10            | India  | -   | -  | -  | - | -   | -   | -           | +   | +      | -   | -     | -     | -     | -     | E       |
| 1994 | MJ-1236         | Bangladesh | - | -  | -  | - | -   | -   | -           | ICE VchBan9 | +   | +      | +   | +     | -     | -     | -     | D       |
| 2002 | CIRS101         | India  | -   | -  | -  | - | -   | -   | ICE VchIndS | + (tr) | +   | -      | -   | -     | -     | -     | -     | B       |
| 2012 | 5 Nampula       | R I R R R S R S I | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 3 Cabo Delgado  | R I R I S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 1 Cabo Delgado  | R S R I S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 1 Nampula       | R I R I S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 1 Nampula       | R I R I S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 1 Cabo Delgado  | R S R S S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 1 Cabo Delgado  | R S R S S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 5 Cabo Delgado  | R R R R S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 3 Cabo Delgado  | R I R I S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 5 Cabo Delgado  | R I R I S R S S | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 3 Niassa        | R R R R S R S S | -           | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B + K   |
| 2013 | 1 Zambézia      | R I R I S R S S | -           | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B       |
| 2013 | 2 Zambézia      | R R R R R S S S | -           | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B       |
| 2014 | 17 Nampula      | R R R R S S S S | -           | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B       |
| 2014 | 11 Sofala       | R R R R S S S S | -           | ICE VchIndS | + (tr) | +   | +      | -   | -           | -   | -     | -     | -     | -     | -     | -     | B       |

https://doi.org/10.1371/journal.pone.0181496.t002
Resistance to Nalidixic acid for tested isolates in the study was 100%, being consistent with data obtained from Iran [21], Haiti [19], Mozambique South rural area [4], and an increasing resistance in Calcutta [15,17]. Conversely, in East Deli [16] and urban South Mozambique [6] was found susceptibility to Nalidixic acid in tested isolates.

In our study, isolates were 100% resistant to Chloramphenicol in 2012, registered a sudden drop of nearly 50% in 2013 with subsequent increase; this behavior was seen in Calcutta [15] and Zambia [11]; while susceptibility to Chloramphenicol was found in East Delhi [16]; and Mozambique [4,6].

Isolates from 2015 were 95% Resistant to Cotrimoxazole, like in Mozambique in 2007 [6], Calcutta [15], Zambia [11], Iran [21], Ghana [17] and East Delhi [16].

Regarding Doxycycline and Azythromycin, recomended drugs for Cholera control by WHO, like in Zambia [11] and Ghana [17], there was some resistance for Doxycyclin and like in urban and rural Bangladesh [12] and Haiti [23], 2012–2013 isolates were susceptible to Azythromycin, while 2015’s like Ghana [17] presented with some resistance.

PCR screening revealed the same virulence genes found in Mozambique by Folgosa [5] and [24], Cholera toxin ctxA and toxin-coregulated pilus, tcpA, contained by the majority of V. cholerae O1 strains, confirming the profile found in V. cholerae O1 El Tor variants B33 and CIRS 101 (CIRS 101 from Bangladesh). Diferent mobilome profiles have been reported for other African countries where V cholerae occurs and unique Mozambique’s profile may be associated with different antibiotic resistance profile.

Due to the plasticity of V. cholerae resulting in the constant emergence of variants, surveillance and characterization of outbreak strains, and their antibiotic resistance determinants, is essential on defining the complex scenario of cholera in this continent as well as worldwide [14].

These findings may be somewhat limited since AST was done according to laboratory discs availability at the moment, usual outbreak scenario where not all antibiotics are tested for each isolate; that could be different on an established antibiotics resistance surveillance scenario, where you predict your demand and therefore your supply. This disabled not only the evaluation of susceptibility profiles for different affected areas and testing for all affected areas but the evaluation of the resistance pattern throughout studied years for Cotrimoxazole, Nitrofurantoin, Azithromycin and Doxycyclin, last two antibiotics specially important because of their reported antimicrobial activity on V. cholerae in previous years and currently in other countries where this pathogen occurs.

Although, serological testing and PCR analyses shows for all strains the same serotype and same genetic similarity, a genetic profile coming from the same clonal origin, it is possible, therefore, to assume a similar resistance pattern for each year.

**Conclusion**

In general, therefore, it seems that antibiotic resistance profile of V. cholerae regarding same serotype and same year varies in different countries and this study strengthens the importance of having local antibiotic choice based on an updated AST local report.

With a highly frequent and increasing resistance, the current data highlights the importance to monitor antimicrobial resistance in epidemic strains, since the appearance of antimicrobial resistance to commonly used and recommended antibiotics will influence Cholera national control strategies.

**Acknowledgments**

Special thanks and gratitude to the Africhol project (Africa network for cholera surveillance) for the financial support; the Mozambican national reference laboratory of Microbiology team...
as well as Provincial field teams, Cláudio Muianga and Dr. Sérgio Chicumbe are acknowledged for their technical support to this study and on writing this manuscript.

**Author Contributions**

**Conceptualization:** Jucunu Elias Chitio, José Paulo Langa.

**Data curation:** Liliana Candida Dengo-Baloi, Dorteia Luisa Inguane.

**Formal analysis:** Liliana Candida Dengo-Baloi, Lena Vania Manhique, Jucunu Elias Chitio, Dorteia Luisa Inguane, José Paulo Langa.

**Funding acquisition:** Cynthia Amino Sema-Baltazar.

**Investigation:** Liliana Candida Dengo-Baloi, Lena Vania Manhique, Jucunu Elias Chitio, Dorteia Luisa Inguane, José Paulo Langa.

**Methodology:** Liliana Candida Dengo-Baloi, Lena Vania Manhique, Jucunu Elias Chitio, Dorteia Luisa Inguane, José Paulo Langa.

**Project administration:** Liliana Candida Dengo-Baloi, Cynthia Amino Semá-Baltazar, Dorteia Luisa Inguane.

**Software:** Dorteia Luisa Inguane.

**Supervision:** Liliana Candida Dengo-Baloi, Cynthia Amino Semá-Baltazar, José Paulo Langa.

**Validation:** Jucunu Elias Chitio, José Paulo Langa.

**Visualization:** Liliana Candida Dengo-Baloi, José Paulo Langa.

**Writing – original draft:** Liliana Candida Dengo-Baloi.

**Writing – review & editing:** Liliana Candida Dengo-Baloi, Cynthia Amino Semá-Baltazar, Lena Vania Manhique, Jucunu Elias Chitio, Dorteia Luisa Inguane, José Paulo Langa.

**References**

1. Sack DA, Organization WH, others. Antimicrobial resistance in shigellosis, cholera, and campylobacteriosis [Internet]. World Health Organization Geneva; 2001 [cited 2015 Aug 3]. Available from: http://cdrwww.who.int/entity/drugresistance/Antimicrobial_resistance_in_shigellosis_cholera_and_cam.pdf
2. Kitaoka M, Miyata ST, Unterweger D, Pukatzki S. Antibiotic resistance mechanisms of Vibrio cholerae. J Med Microbiol. 2011 Apr 1; 60(4):397–407.
3. Aragón M, Barreto A, Tabbard P, Chambule J, Santos C, Nova A. Cholera epidemiology in Mozambique: 1973–1992. Rev Saúde Pública. 1994 Out; 28(5):332–6. PMID: 7660033
4. Mandomando I, Espasa M, Valles X, Sacarai J, Sigauque B, Ruiz J, et al. Antimicrobial resistance of Vibrio cholerae O1 serotype Ogawa isolated in Manhica District Hospital, southern Mozambique. J Antimicrob Chemother. 2007 Jul 23; 60(3):662–4. https://doi.org/10.1093/jac/dkm257 PMID: 17626024
5. Folgosa E, Mastrandrea S, Cappuccinelli P, Uzzau S, Rappelli P, Brian MJ, et al. Molecular identification of pathogenicity genes and ERIC types in Vibrio cholerae O1 epidemic strains from Mozambique. Epidemiol Infect. 2001; 127(01):17–25.
6. Gujral L, Sema C, Rebaudet S, Taibo CLA, Manjate AA, Pierroux R, et al. Cholera Epidemiology in Mozambique Using National Surveillance Data. J Infect Dis. 2013 Nov 1; 208(suppl 1):S107–14.
7. Mengel MA. Cholera in Africa: new momentum in fighting an old problem. Trans R Soc Trop Med Hyg. 2014 Jul 1; 108(7):391–2. https://doi.org/10.1093/trstmh/tru077 PMID: 24836060
8. Greenough WB, Gordon RS Jr, Rosenberg IS, Davies BI, Benenson AS. Tetracycline in the treatment of Cholera. Lancet. 1964; 1:355–7. PMID: 14090856
9. Biemer JJ. Antimicrobial susceptibility testing by the Kirby-Bauer disc diffusion method. Ann Clin Lab Sci. 1973; 3(2):135–40. PMID: 4575155
10. Naha A, Pazhani GP, Ganguly M, Ghosh S, Ramamurthy T, Nandy RK, et al. Development and Evaluation of a PCR Assay for Tracking the Emergence and Dissemination of Haitian Variant ctxB in Vibrio...
cholerae O1 Strains Isolated from Kolkata, India. J Clin Microbiol. 2012 May 1; 50(5):1733–6. https://doi.org/10.1128/JCM.00387-12 PMID: 22357499

11. Mwansa JCL, Mwaba J, Lukwesa C, Bhuiyan NA, Ansaruzzaman M, Ramamurthy T, et al. Multiply antibiotic-resistant Vibrio cholerae O1 biotype El Tor strains emerge during cholera outbreaks in Zambia. Epidemiol Infect. 2007 Jul; 135(05):847.

12. Klontz EH, Das SK, Ahmed D, Ahmed S, Chisti MJ, Malek MA, et al. Long-Term Comparison of Antibiotic Resistances in Vibrio cholerae O1 and Shigella spp. Between Urban and Rural Bangladesh. Clin Infect Dis. 2014;ciu040.

13. Miwanda B, Moore S, Muyembe J-J, Nguefack-Tsagie G, Kabangwa IK, Ndjakani DY, et al. Antimicrobial Drug Resistance of Vibrio cholerae O1 biotype El Tor in Zambia. Vibrio cholerae O1 biotype El Tor strains emerge during cholera outbreaks in Zambia. Emerg Infect Dis. 2007 Jul; 135(05):847.

14. Marin MA, Thompson CC, Fonseca EL, Aboderin AO, Zailani SB, et al. Cholera Outbreaks in Nigeria Are Associated with Multidrug Resistant Atypical El Tor and Non-O1/Non-O139 Vibrio cholerae. Vinetz JM, editor. PLoS Negl Trop Dis. 2013 Feb 14; 7(2):e2049. https://doi.org/10.1371/journal.pntd.0002049 PMID: 2349673

15. Garg P, Chakraborty S, Basu I, Datta S, Rajendran K, Bhattacharya T, et al. Expanding multiple antibiotic resistance among clinical strains of Vibrio cholerae isolated from 1992–7 in Calcutta, India. Epidemiol Infect. 2000; 124(03):393–9.

16. Das S, Choudhry S, Saha R, Ramachandran VG, Kaur K, Sarkar BL. Emergence of multiple drug resistance Vibrio cholerae O1 in East Delhi. J Infect Dev Ctries. 2011; 5(04):294–8.

17. Kuma GK, Opintan JA, Sackey S, Nyarko KM, Opare D, Aryee E, et al. Antibiotic resistance patterns amongst clinical Vibrio cholerae O1 isolates from Accra, Ghana. Int J Infect Control [Internet]. 2014 [cited 2015 Aug 3]; 10(3). Available from: http://www.researchgate.net/profile/Japheth_Opintan/publication/267391039_Antibiotic_resistance_patterns_amongst_clinical_Vibrio_cholerae_O1_isolates_from_Accra_Ghana/links/544e3a430c5f29473161a58cf.pdf

18. Mandal J, Dinoop KP, Parija SC. Increasing antimicrobial resistance of Vibrio cholerae O1 biotype El Tor strains isolated in a tertiary-care centre in India. J Health Popul Nutr. 2012; 12–6. PMID: 22524114

19. Sjolund-Karlsson M, Reimer A, Bolster J, Walker M, Dahouroh G, Batra D, et al. Drug-Resistance Mechanisms in Vibrio cholerae O1 Outbreak Strain, Haiti, 2010. Emerg Infect Dis [Internet]. 2011 Nov [cited 2015 Aug 3]; 17(11). Available from: http://wwwnc.cdc.gov/eid/article/17/11/11-0720_article.htm

20. Mala E, Oberoi A, Alexander VS. Vibrio isolates from cases of acute diarrhea and their antimicrobial susceptibility pattern in a tertiary care hospital. Int J Basic Appl Sci [Internet]. 2013 Dec 6 [cited 2015 Aug 3]; 3(1). Available from: http://www.sciencedepubco.com/index.php/ijbas/article/view/1735

21. Salimi-Khorashad A, Tabatabaee SM, Amirabadi A, Roudbar-Mohamadi S. Vibrio cholerae and Changing of Microbial Resistance Patterns in Sistan and Balouchestan Province. Zahedan J Res Med Sci. 2012; 14(6):63–6.

22. Waturangi DE, Wennars M, Suhartono MX, Wiljaya YF. Edible ice in Jakarta, Indonesia, is contaminated with multidrug-resistant Vibrio cholerae with virulence potential. J Med Microbiol. 2013 Mar 1; 62 (Pt 3):352–9. https://doi.org/10.1099/jmm.0.048769-0 PMID: 23264457

23. Sjolund-Karlsson M, Reimer A, Bolster J, Walker M, Dahouroh G, Batra D, et al. Drug-Resistance Mechanisms in Vibrio cholerae O1 Outbreak Strain, Haiti, 2010. Emerg Infect Dis [Internet]. 2011 Nov [cited 2016 Aug 5]; 17(11). Available from: http://wwwnc.cdc.gov/eid/article/17/11-0720_article.htm

24. Ansaruzzaman M, Bhuiyan NA, Balakrish Nair G, Sack DA, Lucas M, Deen JL, et al. Cholera in Mozambique, Variant of Vibrio cholerae. Emerg Infect Dis. 2004 Nov; 10(11):2057–9. https://doi.org/10.3201/eid1011.040682 PMID: 16010751