Meta-analysis of proportion estimates of Extended-Spectrum-Beta-Lactamase-producing Enterobacteriaceae in East Africa hospitals

Tolbert Sonda1,2*, Happiness Kumburu1,2 , Marco van Zwetselaar1, Michael Alifrangis3,4, Ole Lund6, Gibson Kibiki1,2 and Frank M. Aarestrup5

Abstract

Background: A high proportion of Extended-Spectrum-Beta-Lactamase (ESBL) producing Enterobacteriaceae is causing common infections in all regions of the world. The burden of antibiotic resistance due to ESBL in East Africa is large but information is scarce and thus it is unclear how big the problem really is. To gain insight into the magnitude and molecular epidemiology of ESBL-producing Enterobacteriaceae in East Africa a literature search was performed in PubMed on 31 July 2015 to retrieve articles with relevant information on ESBL.

Methods and results: Meta-analysis was performed to determine overall proportion estimate of ESBL-producing Enterobacteriaceae. A total of 4076 bacterial isolates were included in the analysis. The overall pooled proportion of ESBL-producing Enterobacteriaceae among included surveys done in East African hospitals was found to be 0.42 (95 % CI: 0.34–0.50). Heterogeneity (I²) between countries’ proportions in ESBL was significantly high (96.95 % and p < 0.001). The frequently detected genes encoding ESBL were CTX-M, TEM, SHV and OXA while the most infrequent reported genes were KPC and NDM.

Conclusion: The available studies show a very wide variation in resistance due to ESBL between countries. This highlights a need for active surveillance systems which can help understand the actual epidemiology of ESBL, aid in formulating national or regional guidelines for proper screening of ESBL, and support developing standardized approaches for managing patients colonized with ESBL.

Keywords: Antibiotic resistance, Extended-Spectrum-Beta-Lactamase, ESBL, Enterobacteriaceae, East Africa

Background

The production of beta-lactamases is the most common mechanism for bacteria to acquire resistance to broad-spectrum beta-lactam antibiotics. These hydrolytic enzymes are encoded by various gene variants. TEM, named after Temoneira, is one of the first enzymes identified in Europe in the 1960s in Escherichia coli [1]. Since then, several other enzymes like CTX, SHV, OXA have been reported in different parts of the world [2–8]. Extended-Spectrum-Beta-Lactamase (ESBL) producing Enterobacteriaceae can be defined as those producing β-lactamases capable of conferring bacterial resistance to the penicillins, first-, second-, and third-generation cephalosporins, and aztreonam (but not the cephamycins or carbapenems) by hydrolysis of these antibiotics, and which are inhibited by β-lactamase inhibitors such as clavulanic acid [9].

The proportion of ESBL-producing bacteria causing common infections in all regions of the world is high, making antibiotic resistance due to ESBL being a major global public health problem [10]. Patients infected with ESBL not only have an increased risk of treatment failure, sometimes resulting in death, but also require more health-care resources. ESBL bacterial infections are...
becoming challenging because physicians run out of drug options. Although there might be differences in magnitude depending on region or country, ESBLs used to be considered primarily nosocomial. Currently they can be frequently found in both hospitals and communities, though magnitudes reported in community-based surveys are generally lower [4, 6–8, 11–13]. Several risk factors have been documented to be associated with ESBL acquisition, including: previous hospitalisation, previous use of antibiotics such as third generation cephalosporins, hospital overcrowding, bed sharing when hospitalised, and international travel [3, 14–19].

Due to regional differences in the ESBL proportion and distribution, therapeutic decisions should be based on local guidelines derived from local epidemiological data [20]. In resource-rich countries many surveillance systems have been set up to estimate the burden of bacterial infections including ESBL and to determine risk factors for acquisition of ESBL bacteria as well as the clinical outcomes associated with their infection [10, 21–23]. In East Africa however, only scarce and scattered information is available on ESBL epidemiology and risk factors associated with ESBL bacterial infection. When available, the data mostly originates from hospital-based studies [7, 24–28]. To gain a better insight in the proportion estimates and molecular epidemiology of ESBL-producing Enterobacteriaceae in East Africa, we retrieved available peer-reviewed articles and collated the information in this review article. The PRISMA statement was used to guide the meta-analysis [29]; the PRISMA checklist is available as supplemental material (Additional file 1).

Materials and methods

Literature search and selection

We defined the East African region as the countries forming the East African Community (Burundi, Kenya, Rwanda, Tanzania and Uganda), with the addition of Ethiopia, based on the UN geographical regions definition. A PubMed search was performed on 31 July 2015 using the search query “(Burundi OR Kenya OR Rwanda OR Tanzania OR Uganda OR Ethiopia) AND (ESBL OR extended-spectrum-beta-lactamase).” A total of 29 potential articles were found. Articles studying non-human subjects, review articles, and articles describing isolates from outside the six countries mentioned above were excluded. A total of 24 articles were included (Fig. 1).

Data extraction and analysis

For data consistency, two people extracted data independently from each article. Whenever there was discordance in the data extracted, consensus was reached by double-checking the article. Data extracted included: name of first authors, year of study (or publication if year of study was not documented), department, target population, isolate source and common species isolated. Other data extracted were the methods used to test for ESBL, the number of Enterobacteriaceae analysed, the number of Enterobacteriaceae positive for ESBL, the gene variants encoding for ESBL, and risk factors for ESBL bacterial infection.

Stata version 13.1 (College Station, Texas 77845 USA) was used to perform meta-analysis of the proportion of ESBL-producing Enterobacteriaceae as described by Nyaga et al. [30]. In the analysis, a random-effects meta-analysis model was used to calculate the pooled (weighted) proportion of ESBL and the I² statistic (measure of inconsistency). The I² statistic expresses the percentage of total variation across studies due to heterogeneity. A value of 0 % shows no observed heterogeneity, increasing values indicate increasing heterogeneity.

Results

Distribution of articles describing ESBL in East Africa

The 24 articles reviewed were from cross-sectional hospital-based studies. 4 (16.7 %) were from Ethiopia, 4 (16.7 %) were from Kenya, 12 (50 %) were from Tanzania, 3 (12.5 %) were from Uganda, and 1 (4.2 %) was from Rwanda. No articles were found for Burundi. 18 (75 %) of reviewed articles included patients attending in-patient and out-patient departments, 4 (16.7 %) included patients attending in-patient departments only, and 2 (8.3 %) included patients attending out-patient departments only. 20 (83.3 %) of the reviewed articles studied E. coli plus other species while 19 (79.2 %) studied K. pneumoniae plus other species.

One (4.2 %) article (from Rwanda) investigated the risk factors for infection with ESBL-producing Enterobacteriaceae. In this article, previous use of cephalosporins, ciprofloxacin and hospitalisation were found to be significant risk factors for ESBL bacterial infection.

Laboratory methods used to estimate the proportion of ESBL

Sixteen (66.7 %) of the articles reviewed used Double Disk Synergy Test (DDST) alone; 2 (8.3 %) articles used PCR-sequencing alone while 6 (25 %) articles used both DDST and PCR-sequencing methods to estimate ESBL proportions.

Proportion estimates of ESBL in East Africa

Based on the available data (Fig. 2), East Africa’s overall pooled proportion of ESBL-producing Enterobacteriaceae was 0.42 (95 % CI: 0.34–0.50). Overall heterogeneity was significant (I² 95.95 %, p < 0.001). The pooled proportion of ESBL-producing Enterobacteriaceae in Ethiopia was 0.30 (95 % CI: 0.21–0.38), I² was 67.98 % and p = 0.02. The pooled proportion of ESBL-producing Enterobacteriaceae in Kenya was 0.47 (95 % CI: 0.23–0.71); I² was
98.82 % and \( p < 0.001 \). The pooled proportion of ESBL-producing *Enterobacteriaceae* in Tanzania was 0.39 (95 % CI: 0.30–0.48); \( I^2 \) was 93.16 % and \( p < 0.001 \). The pooled proportion of ESBL-producing *Enterobacteriaceae* in Uganda was 0.62 (95 % CI: 0.38–0.87); \( I^2 \) was 97.83 % and \( p < 0.001 \).

The intra-country heterogeneity is reflected in the range of the estimates of the ESBL proportion. In Tanzania, the median (range) ESBL proportion estimate was 38.8 % (14.2–75.9). In Kenya the median ESBL proportion was 45.8 % (13.1–88.3). Ethiopia recorded median ESBL proportion of 30.9 % (21.9–40.4). In Uganda the recorded median ESBL proportion was 61.7 % (43.8–81.4). The single article for Rwanda reported a 38.3 % ESBL proportion (Fig. 3).

**Molecular epidemiology of ESBL in East Africa**

Out of 24 reviewed articles, 8 (33.3 %) of which, 4 (50 %) from Tanzania, 3 (37.5 %) from Kenya and 1 (12.5 %) from Uganda had data on ESBL-encoding genes. The predominant ESBL-encoding genes reported were CTX-M, TEM and SHV. Five articles provided proportion estimates for these genes (Table 1). One article reported no quantitative data but identified the same three genes [12]. One article identified CTX-M-15, CMY-2 and AmpC [31]. One article reported the multi resistant and rarer genes VIM (12.3 %), OXA-48 (4.9 %), KPC (3.5 %) and NDM (3.1 %) genes [42].

**Discussion**

The scarcity of studies available from the East African region warrants caution in drawing conclusions. Little information overall is available, with no studies on Burundi and only one on Rwanda. However, this review finds high proportion estimates of ESBL-producing *Enterobacteriaceae* across hospitals in the East African region.

The overall pooled ESBL proportion estimate for East African hospitals (42 %) is close to estimates for Ghana (49 %), Cameroon (54 %), Gabon (45 %) and Morocco (43 %) [18, 32–34]. This estimate is also close to data reported for China, where a nationwide survey that included 30 hospitals reported over 46 % resistance due to ESBL [35]. However the East African proportion is considerably higher than averages reported for resource-rich countries. For instance, the average ESBL proportion estimates reported in a nationwide hospital survey in Germany for 2012 were in the 10 to 15 % range [36]. In 2012, a study for nine US census regions reported ESBL proportion estimates in the 4 to 12 % range [37], while a 9-year survey in Japan recorded a proportion estimate of 6.3 % in 2003 increasing to 10–20 % in 2011 [38]. It should be noted that this study focused on community-acquired infections whereas the other studies concerned nosocomial-infections.

In this review, inter-country and inter-study results show a wide and statistically significant degree of variation.
in proportion estimates \((p < 0.05 \text{ in all cases})\). There are several possible factors that may account for the variations seen in this review. The first factor is the difference in sensitivity and specificity between methods used in estimating proportions. Some studies reviewed estimated ESBL proportions using purely phenotypic methods, while others used both phenotypic and molecular-based methods. The study done in Uganda shows the proportion estimate using genotypic methods being higher (44\%) than when the same isolates were screened for ESBL using phenotypic methods (18\%) [39].

A second factor contributing to the variation in proportion estimates is type of wards or units, site of infection, type of specimen collected and whether patients

| Study | ES (95% CI) |
|-------|-------------|
| Ethiopia |            |
| Seid et al (2004) | 0.40 (0.28, 0.54) |
| Beyene et al (2006) | 0.26 (0.17, 0.37) |
| Mululem et al (2012) | 0.36 (0.24, 0.48) |
| Esthete et al (2014) | 0.22 (0.16, 0.29) |
| Subtotal \((I^2 = 67.98\%, p = 0.02)\) | 0.30 (0.21, 0.38) |
| Kenya |            |
| Kariuki et al (2007) | 0.65 (0.38, 0.86) |
| Kohli et al (2008) | 0.14 (0.11, 0.18) |
| Kiru et al (2010) | 0.27 (0.24, 0.30) |
| Maina et al (2012) | 0.88 (0.77, 0.96) |
| Subtotal \((I^2 = 98.62\%, p = 0.00)\) | 0.47 (0.23, 0.71) |
| Rwanda |            |
| Muvunyi et al (2009) | 0.38 (0.31, 0.45) |
| Tanzania |            |
| Blomberg et al (2001) | 0.14 (0.09, 0.22) |
| Ndugulilo et al (2003) | 0.26 (0.13, 0.42) |
| Mushiri et al (2007) | 0.35 (0.29, 0.41) |
| Kayanga et al (2009) | 0.47 (0.35, 0.59) |
| Meshana et al (2009) | 0.29 (0.25, 0.34) |
| Meshana et al (2009) | 0.50 (0.43, 0.58) |
| Moyo et al (2010) | 0.45 (0.39, 0.51) |
| Mawalla et al (2011) | 0.67 (0.45, 0.84) |
| Moremi et al (2012) | 0.34 (0.26, 0.44) |
| Manyah et al (2012) | 0.76 (0.56, 0.90) |
| Nsik et al (2013) | 0.14 (0.08, 0.22) |
| Ahmed et al (2013) | 0.43 (0.32, 0.54) |
| Subtotal \((I^2 = 93.16\%, p = 0.00)\) | 0.39 (0.30, 0.48) |
| Uganda |            |
| Sani et al (2011) | 0.81 (0.74, 0.87) |
| Moses et al (2012) | 0.44 (0.39, 0.49) |
| Katerega et al (2014) | 0.62 (0.52, 0.71) |
| Subtotal \((I^2 = 97.63\%, p = 0.00)\) | 0.62 (0.36, 0.87) |

Heterogeneity between groups: \(p = 0.000\)

Overall: \((I^2 = 98.95\%, p = 0.00)\); 0.42 (0.34, 0.50)
were attending out-patient or in-patient departments. Hospitalised patients especially in intensive care units are generally at a higher risk of acquiring nosocomial infections, which are likely to be ESBL-producers than patients attending out-patient departments [6, 7, 40–42]. The Rwanda study [15] reports ESBL proportions of 38 % and 5.9 % among inpatients and outpatients respectively, within a single hospital. Similar findings have been documented in Cameroon where ESBL proportions were 23.1 and 6.7 % among outpatients and healthy volunteers respectively [16].

Many reports have documented the difference in ESBL proportion estimates between hospitals and congested centers (such as orphanages) versus community-based surveys [15, 16, 43–46]. The lack of any estimates for community-based ESBL carriage in East Africa underscores an urgent need for surveillance in the region. Infection control in hospitals including hand hygiene and rational antibiotic use can be effective measures to stop further spread of the ESBL-producing Enterobacteriaceae in both hospitals and communities.

We noted that few (33.3 %) articles investigated different genes encoding ESBL [6, 12, 24, 31, 39, 47–49]. The most common gene variants in these articles are those encoding CTX, TEM and SHV. However one study in Tanzania reported occurrence of genes such as NDM that confer resistance to Carbapenem [47]. With such little information in hand, it is difficult to devise focused and effective interventions for containing resistance. Data from other parts of the world, notably from resource-rich countries, might not be generalisable to African settings.

Risk factors associated with resistance due to ESBL are key for planning the optimal approach to managing the problem. Of the reviewed articles only the one study done in Rwanda investigated these factors. It was concluded that previous use of ciprofloxacin, third-generation cephalosporin and being an inpatient or hospitalised are risk factors for ESBL carriage [15]. Similarly, several studies done in other sub-Saharan African countries have documented the previous use of antibiotics (ciprofloxacin), previous hospitalisation, overcrowding in hospitals, and bed sharing to be associated with ESBL colonisation [14, 16, 17]. These findings are consistent with several studies done outside of Africa [50–58].

In African settings previous antibiotics use is not entirely dependent on previous hospitalisation. Over-the-counter sale of drugs or self-medication, consumption of counterfeit drugs, improper dosage and non-adherence are very common. These practices unnecessarily fuel the process of positive selection for many types of antibiotic resistance. The majority of sub-Saharan hospitals lack antibiotics resistance monitoring systems, proper antibiotic usage guidelines and proper hospital premises disinfection guidelines. If we do not put strong pragmatic measures in place our hospitals may soon turn into hotspots not only for ESBL-producing Enterobacteriaceae, but also for many other types of resistant microorganisms.

When interpreting the data compiled in this review, a number of limitations must be taken into account. All studies took place in university teaching or referral hospitals, which will have a different case-mix from peripheral health centres, and will generally be located in urban areas. Another source of bias is the fact that no study included community-based findings. As in any meta-analysis, the pooled proportions across studies must be interpreted with care, as protocols for observational studies are not
| Country     | Year | Hospital       | Department   | Population      | Specimen                  | Method               | Species | Isolates (N) | ESBL (%) | ESBL-Genes  |
|-------------|------|----------------|--------------|-----------------|---------------------------|----------------------|---------|--------------|-----------|-------------|
| Ethiopia    | 2004 | Regional       | Both         | Children, adults | Urine, pus, sputum       | DDST                 | K. pneumoniae | 57          | 42         | ND          |
| Ethiopia    | 2014 | University     | Both         | Children, adults | Urine                    | DDST                 | K. pneumoniae, E. coli | 183        | 22         | ND          |
| Ethiopia    | 2006 | University     | Out-patient  | Children        | Blood, stool              | DDST                 | Salmonella spp     | 81          | 27         | ND          |
| Ethiopia    | 2012 | University     | Both         | Children, adults | Urine, pus, sputum, stool | DDST                 | E. coli          | 67          | 36         | ND          |
| Kenya       | 2010 | Not Stated     | Both         | Children, adults | Urine, stool, blood     | DDST, PCR-Sequencing  | E. coli          | 912         | 27         | CTX-M (78 %), SHV (3–5 %), TEM (16 %) |
| Kenya       | 2007 | University     | Both         | Children, adults | Urine                    | PCR-Sequencing       | E. coli          | 17          | 71         | CTX-M-15 type ESBLs and CMY-2 AmpC |
| Kenya       | 2008 | University     | Both         | Children, adults | Blood                    | DDST                 | K. pneumoniae, E. coli | 359        | 14         | ND          |
| Kenya       | 2012 | University     | Both         | Children, adults | Urine, pus, sputum, stool | DDST, PCR-Sequencing  | K. pneumoniae, E. coli | 52          | 89         | CTX-M (88.5 %), blaSHV (25 %), TEM (34.6 %) |
| Rwanda      | 2009 | University     | Both         | Children, adults | Urine                    | DDST                 | K. pneumoniae, E. coli | 196        | 38         | ND          |
| Tanzania    | 2001 | University     | In-patients  | Children        | Blood                    | DDST, PCR-Sequencing  | K. pneumoniae, E. coli | 125        | 15         | CTX-M, SHV, TEM |
| Tanzania    | 2009 | University     | In-patients  | Children, adults | Urine, pus, wound, blood | DDST, PCR-Sequencing  | K. pneumoniae     | 183        | 50         | CTX-M-15 (76 %), TEM-104 (19 %), TEM-176 (2 %), SHV-11 (3.2 %) |
| Tanzania    | 2003 | University     | In-patients  | Children, adults | Urine, wound, blood      | DDST, PCR-Sequencing  | E. coli, Enterobacter spp, others | 39          | 28         | TEM (55 %), SHV (64 %), CTX-M (45.4 %) |
| Tanzania    | 2010 | University     | Both         | Children, adults | Urine                    | DDST                 | K. pneumoniae, E. coli | 270        | 45         | ND          |
| Tanzania    | 2009 | University     | Both         | Children, adults | Urine, pus, wound, blood | DDST                 | K. pneumoniae, Escherichia coli, Acinetobacter spp | 377        | 29         | ND          |
| Tanzania    | 2013 | University     | In-patients  | Women, neonates  | Rectal swabs             | DDST                 | K. pneumoniae, Enterobacter spp | 113        | 15         | ND          |
| Tanzania    | 2013 | University     | Both         | Children        | Urine                    | DDST                 | K. pneumoniae, E. coli, others | 84          | 44         | ND          |
| Tanzania    | 2012 | University     | Both         | Children, adults | Pus                      | DDST                 | K. pneumoniae, E. coli, others | 29          | 79         | ND          |
| Tanzania    | 2007 | University     | Both         | Children, adults | Urine, pus, blood        | PCR-Sequencing       | K. pneumoniae, E. coli, P. aeruginosa, others | 227        | 35         | VIM (12.3 %), OXA-48 (4.9 %), KPC (3.5 %), NDM (3.1 %) |
| Tanzania    | 2012 | University     | Both         | Children, adults | Wound                    | DDST                 | Pseudomonas spp, Proteus spp, K. pneumoniae, E. coli | 117        | 35         | ND          |
| Tanzania    | 2011 | University     | Both         | Children, adults | Wound                    | DDST                 | K. pneumoniae, E. coli, others | 24          | 71         | ND          |
| Tanzania    | 2009 | University     | Both         | Children        | Blood                    | DDST                 | K. pneumoniae, E. coli, others | 72          | 49         | ND          |
Table 1 Distribution of articles reviewed on resistance due to ESBL in East Africa hospitals and common gene variants encoding for ESBL (Continued)

| Country [Ref] | Year | Region | Type | Site(s) | Method(s) | Genotypes | Zn | Interpretation |
|---------------|------|--------|------|---------|-----------|-----------|----|----------------|
| Uganda [10]   | 2011 | Regional | Both | Children, adults | Wound | DDST | K. pneumoniae, E. coli, others | 145 | 81 | ND |
| Uganda [60]   | 2014 | Regional | Both | Children, adults | Urine, blood, wound, CSF* | DDST | E. coli, K. pneumoniae, P. mirabilis, others | 115 | 62 | ND |
| Uganda [31]   | 2012 | Regional | Both | Children, adults | Urine, pus, wound, sputum, stool, CSF, vaginal-swabs | DDST, PCR-Sequencing | K. pneumoniae, E. coli, Salmonella spp, others | 484 | 44 | CTX-M (70 %), SHV (34 %) and TEM (47 %) |

*Both, out-and in-patients departments
*DDST double disc synergy test
*ND not determined
*PCR polymerase chain reaction
*CSF cerebrospinal fluid
standardised across the studies included in the review. A notable factor contributing to variation is the different mix of in- and outpatients between the studies. Finally, the authors acknowledge that more data on ESBL in East Africa may be available from sources other than those searched for this review.

Conclusion
The burden of antibiotic resistance due to ESBL is present across East African region. Little information overall is available, and close to none for Burundi. The available studies present proportion estimates due to ESBL with a wide degree of variation. The scarcity of data on predictors, clinical outcomes, magnitudes and gene variants encoding resistance due to ESBL-producing Enterobacteriaceae calls for active surveillance systems, which can help understand the current epidemiology of ESBL within the region. Furthermore this can aid in developing national and regional guidelines for proper screening of ESBL as well as developing standardized approaches for managing patients colonized with ESBL-producing Enterobacteriaceae.

Additional file

Additional file 1: The PRISMA checklist. (DOC 62 kb)

Competing interests
The authors declare that they have no competing interests.

Authors' contributions
TS and GK conceived the idea. TS and MZ gathered data, analyzed them and prepared manuscript draft. All authors read, revised and approved the final manuscript.

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Author details
1 Kilimanjaro Clinical Research Institute, Kilimanjaro Christian Medical Centre, Moshi, Tanzania. 2 Kilimanjaro Christian Medical University College, Moshi, Tanzania. 3 Centre for Medical Parasitology, Department of Immunology and Microbiology, University of Copenhagen, Copenhagen, Denmark. 4 Department of Infectious Diseases, Copenhagen University Hospital, Copenhagen, Denmark. 5 Centre for Genomic Epidemiology, Technical University of Denmark, Copenhagen, Denmark. 6 Centre for Biological Sequence Analysis, Technical University of Denmark, Copenhagen, Denmark.

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