RESEARCH ARTICLE

Sepsis Caused by Extended-Spectrum Beta-Lactamase (ESBL)-Positive *K. pneumoniae* and *E. coli*: Comparison of Severity of Sepsis, Delay of Anti-Infective Therapy and ESBL Genotype

Christian Sakellariou1*, Stephan Gürntke1*, Ivo Steinmetz2, Christian Kohler2, Yvonne Pfeifer3, Petra Gastmeier1, Frank Schwab1, Axel Kola1, Maria Deja4, Rasmus Leistner1*

1 Institute of Hygiene and Environmental Medicine, National Reference Center for the Surveillance of Nosocomial Infections, Charité Universitätsmedizin Berlin, Hindenburgdamm 27, 12203 Berlin, Germany, 2 Friedrich Löffler Institute of Medical Microbiology, Universitätsmedizin Greifswald, Martin-Luther-Str.6, 17475, Greifswald, Germany, 3 Robert Koch Institute, FG13 Nosocomial Pathogens and Antibiotic Resistance, 38855, Wernigerode, Germany, 4 Department of Anesthesiology and Intensive Care, Charité Universitätsmedizin Berlin, Campus Benjamin Franklin, Hindenburgdamm 30, 12203, Berlin, Germany

*These authors contributed equally to this work.

* rasmus.leistner@charite.de

Abstract

Infections with extended-spectrum beta-lactamase-producing *Enterobacteriaceae* (ESBL-E) are associated with increased mortality. Outcome differences due to various species of ESBL-E or ESBL genotypes are not well investigated. We conducted a cohort study to assess risk factors for mortality in cases of ESBL-E bacteremia (*K. pneumoniae* or *E. coli*) and the risk factors for sepsis with organ failure. All consecutive patients of our institution from 2008 to 2011 with bacteremia due to ESBL-E were included. Basic epidemiological data, underlying comorbidities, origin of bacteremia, severity of sepsis and delay of appropriate anti-infective treatment were collected. Isolates were PCR-screened for the presence of ESBL genes and plasmid-mediated AmpC β-lactamases. Cox proportional hazard regression on mortality and multivariable logistic regression on risk factors for sepsis with organ failure were conducted. 219 cases were included in the analysis: 73.1% due to *E. coli*, 26.9% due to *K. pneumoniae*. There was no significant difference in hospital mortality (ESBL-*E. coli*, 23.8% vs. ESBL-*K. pneumoniae* 27.1%, *p* = 0.724). However, the risk of sepsis with organ failure was associated in cases of *K. pneumoniae* bacteremia (OR 4.5, *p* < 0.001) and patients with liver disease (OR 3.4, *p* = 0.004) or renal disease (OR 6.8, *p* < 0.001). We found significant differences in clinical presentation of ESBL-E bacteremia due to *K. pneumoniae* compared to *E. coli*. As *K. pneumoniae* cases showed a more serious clinical presentation as *E. coli* cases and were associated with different risk factors, treatment and prevention strategies should be adjusted accordingly.
Introduction

Infections due to extended-spectrum beta-lactamase-producing Enterobacteriaceae (ESBL-E) are associated with impaired outcome compared to infections with susceptible pathogens [1–3]. Former studies on ESBL-E bacteremia proved that a delay of adequate antimicrobial chemotherapy can be an important factor on mortality [1, 2, 4]. This effect seems to be the most evident in cases of septic shock or organ failure [5]. This is even more important since there is evidence that infections due to K. pneumoniae are associated with a worse course compared to other Enterobacteriaceae [4, 6, 7]. However, studies concerning ESBL-E infections often do not differentiate between the infecting species. There are only few studies comparing outcome parameter of different Enterobacteriaceae [4, 6, 8–10]. To analyze the effect of the different ESBL-E species on mortality and on the clinical presentation, we conducted a cohort study comparing cases of ESBL-positive K. pneumoniae- and E. coli bacteremia including data on the timing of their antimicrobial treatment.

Methods

Study design

We conducted a retrospective cohort study on patients with bacteremia due to ESBL-E. The setting of this study was the Charité University Hospital in Berlin, Germany, a tertiary care university hospital with over 120,000 admissions per year. Previously, our ethics committee approved the study without informed consent. The patient data based on secondary clinical information. The biological material was obtained clinically and analyzed by a separate institution after anonymization in our institution (reference number EA4/031/11). Parts of the study have been published in different analyses including differing data sets [10, 11].

All inpatients with ESBL-E bacteremia (E. coli or K. pneumoniae) diagnosed between January 1st, 2008 and December 31st 2011 were included. The bacteremia was classified as hospital onset if a positive blood culture was collected after the third hospital day. In case of more than one cultured organism in the first positive blood culture, the episode was defined as polymicrobial bacteremia. If the patient showed multiple subsequent positive blood cultures, the first positive blood culture lead to the allocation to one of the two pathogens and this episode was included in the analysis. Underlying comorbidities were assessed according to the method by Charlson et al. [12]. The comorbidities were collected using the patients’ ICD-10 codes and grouped for the calculation of the Charlson comorbidity index (CCI) according to the method by Thygesen et al. [13].

Further clinical parameters were collected by analysis of the patients’ files. To assess the origin of ESBL bacteremia we collected data on earlier infections during the analyzed hospital stay. These infections were due to same ESBL-positive organisms as the corresponding bacteremia episode and at maximum 14 days prior to the bacteremia onset. The infections were assessed according to the CDC definitions [14]. Primary bacteremia was defined as central venous catheter (CVC) <48h prior to the bacteremia onset without presence of another ESBL-E infection. Mortality was defined as in-hospital mortality and ESBL colonization was assessed as colonization with ESBL-E prior to the bacteremia episode at any site. Sepsis, severe sepsis, and septic shock were defined according to the definitions of the consensus conference of the American College of Chest Physicians and the Society of Critical Care Medicine which also find application in Germany [15, 16]. Delay of anti-infective treatment (DAT) was defined as initiation of effective treatment ≥1 day after onset of bacteremia. Effective treatment was defined as an antimicrobial agent the ESBL-positive organism was tested susceptible against.
Microbiological methods

Species identification and antimicrobial susceptibility testing was performed using the Vitek 2 automated system with interpretation and antibiogram reporting according to the CLSI standard [17]. Confirmation of ESBL production was performed by a minimum inhibitory concentration dilution test on a multiwell microtiter plate. Three cephalosporins (cefotaxime, ceftazidime, cefpodoxime) were tested alone and in combination with ESBL inhibitor clavulanic acid. All verification swabs were inoculated onto chrome ID ESBL agar (BioMerieux). ESBL-positive isolates were affirmed by Double Disc Synergie Testing using 3rd generation cephalosporins with/without clavulanic acid (Mast). Species confirmation was done by API20E (BioMerieux). The isolates were screened for the presence of different ESBL genes (blaTEM-type, blaSHV-type, blaCTX-M-1/2/9 group) by polymerase chain reaction (PCR) and subsequent sequencing [18]. If none of these ESBL genes could be identified, additional PCR tests for the presence of plasmid-mediated AmpC β-lactamases [19] and further ESBL genes (blaCTX-M-8-type, blaCTX-M-26-type) were performed [20]. Furthermore, basic bacterial typing of all ESBL-positive E. coli isolates was performed by a PCR-based method for determination of the four major E. coli phylogenetic groups [21].

Statistical methods

Parameters in the univariate analysis were tested using the Wilcoxon rank sum test for continuous variables and the Fisher’s exact test for categorical variables. For the analysis of factors associated to severity of illness, the categories were transformed to a binary category as bacteremia (bacteremia and sepsis) vs. sepsis with organ failure (severe sepsis and septic shock). Clinical parameter of E. coli- and K. pneumoniae- cases were compared using univariate analysis. Multivariable binary logistic analysis using a stepwise forward regression was computed for the analysis of clinical risk factors for sepsis with organ failure. We compared the clinical parameter of the deceased patients and the alive discharged patients using univariate analysis. We calculated adjusted hazard ratios for in-hospital death using Cox-proportional hazard regression using a stepwise forward approach. In both multivariable analyses, parameters with p-values <0.20 in the univariate analyses were considered in the analysis. Variables with p-values <0.05 were included and variables with p ≥ 0.05 were excluded. All tests of were two tailed with a p-value <0.05 considered to be significant. Data was analyzed using IBM SPSS Statistics Version 22.

Results

We identified altogether 243 consecutive cases of bacteremia, 177 cases due to ESBL-positive E. coli (73.1%) and 66 cases due to K. pneumoniae (27.2%) together. No case showed both analyzed pathogens in the same blood culture. From twenty-three cases (9.9%), sufficient data was not available and therefore excluded. The remaining 219 cases were analyzed (n = 160, 73.1% due to E. coli and n = 59, 26.9% due to K. pneumoniae). We found an increasing risk for mortality in relation to the applied definitions for severity of sepsis (Fig 1). The characteristics of the analyzed cohort stratified by infecting organism are given in S1 Table. Patients suffering from ESBL-positive K. pneumoniae bacteremia (ESBL-KP-Bac) were younger than the compared patients with ESBL-positive E. coli bacteremia (ESBL-EC-Bac). They more often had a secondary bacteremia due to a surgical site infection and more often an unknown source of infection. ESBL-KP-Bac was more often associated with sepsis with organ failure. Patients with sepsis with organ failure showed significantly reduced DAT compared to patients presenting with bacteremia only (Median 0 days, IQR 0;2 days vs. Median 2 days, IQR 0;3 days, p = 0.003). Patients with ESBL-KP-Bac showed an increased mortality compared to
ESBL-EC-Bac (27.1% vs. 23.8%) but not statistical significant. ESBL-EC-Bac however, was more common among patients with previous urinary tract infection. The mostly used antimicrobial agents after infection onset were carbapenems: 71.2% (n = 156), quinolones 12.3% (n = 27), tigecyclin 5.0% (n = 11), piperacillin-tazobactam 2.3% (n = 5) and gentamicin 2.3% (n = 5).

Microbiology parameter

In the clinically reported antibiogram, 98.6% (n = 216) isolates were resistant to piperacillin/tazobactam, 98.2% (n = 215) to ceftazidime, 69.4% (n = 152) to ciprofloxacin and 43.4% (n = 95) to gentamicin. None of the included isolates was reported resistant to the carbapenems imipenem or meropenem. Of the 219 isolates, 88.5% (n = 194) were available for further ESBL genotype analysis; the remaining 25 were not retrievable. The distribution of the ESBL genotypes is shown in Tables 1 and 2 with overall CTX-M-15, CTX-M-1, CTX-M-14 and SHV-5 as most common types. One-hundred and seven isolates (55.2%) carried two or more TEM- or SHV-type beta-lactamases. Seven isolates (3.6%) did not carry an ESBL gene. Five showed either TEM-181, TEM-1 or SHV-1 overproduction, one was CMY-positive and one did not show any beta-lactamase at all. The distribution pattern of the phylogenetic groups within the analyzed 140 E. coli isolates was B2 (33.6%, n = 47), A (28.6%, n = 40), D (26.4%, n = 37) and B1 (11.4%, n = 16).

Factors associated with sepsis with organ failure

Table 3 shows the results of the multivariable analysis on risk factors for sepsis with organ failure. The K. pneumoniae cases were associated with 4.5 times higher odds for an organ failure at presentation. Furthermore, sepsis with organ failure was associated with liver disease (OR 3.3) and moderate to severe renal disease (OR 6.835).
Table 1. Univariate analysis of clinical parameter in patients presenting with sepsis with organ failure and bacteremia.

| Parameter | Bacteremia (n = 138) | Sepsis with organ failure (n = 81) | P-value |
|-----------|----------------------|----------------------------------|---------|
| Age years / Age < 61 years | 67 (48.6%) | 42 (51.9%) | 0.676 |
| Male sex | 93 (67.4%) | 53 (65.4%) | 0.769 |
| Charlson comorbidity index > 6 | 63 (45.7%) | 51 (63.0%) | 0.017 |
| In-hospital death | 12 (8.7%) | 42 (51.9%) | <0.001 |
| Polymicrobial bacteraemia | 19 (13.8%) | 10 (12.3%) | 0.839 |
| Hospital onset | 74 (53.6%) | 59 (72.8%) | 0.006* |
| ESBL colonization before onset | 85 (61.6%) | 53 (65.4%) | 0.664 |
| Bacteraemia due to *E. coli* | 115 (83.3%) | 45 (55.6%) | <0.001* |
| Bacteraemia due to *K. pneumoniae* | 23 (16.7%) | 36 (44.4%) | |

**Origin of ESBL-E bacteraemia**

| | Bacteremia (n = 138) | Sepsis with organ failure (n = 81) |
|-----------------|----------------------|----------------------------------|
| Urinary tract infection | 55 (39.9%) | 20 (24.7%) | 0.027* |
| Lower respiratory tract infection | 22 (15.9%) | 18 (22.2%) | 0.279 |
| Intra-abdominal infection | 9 (6.5%) | 13 (16.0%) | 0.035* |
| Surgical site infection | 4 (2.9%) | 4 (4.9%) | 0.472 |
| Primary bacteraemia | 19 (13.8%) | 11 (13.6%) | 1.000 |
| Other | 6 (4.3%) | 2 (2.5%) | 0.756 |
| Unknown Origin | 35 (25.4%) | 19 (23.4%) | 0.883 |

**Severity of illness and delay of anti-infective treatment**

| | Bacteremia (n = 138) | Sepsis with organ failure (n = 81) |
|-----------------|----------------------|----------------------------------|
| Delayed adequate anti-infective treatment | 70 (50.7%) | 25 (30.9%) | <0.001 |

**ESBL Genotype**

| | Bacteremia (n = 138) | Sepsis with organ failure (n = 81) |
|-----------------|----------------------|----------------------------------|
| No ESBL genotype | 4 (2.9%) | 3 (3.7%) | 0.711 |
| CTX-M-1 | 30 (21.7%) | 9 (11.1%) | 0.066* |
| CTX-M-14 | 8 (5.8%) | 6 (7.4%) | 0.776 |
| CTX-M-15 | 60 (43.5%) | 38 (46.9%) | 0.674 |
| CTX-M-2 | 1 (0.7%) | - | 1.000 |
| CTX-M-2/97 | 2 (1.4%) | - | 0.532 |
| CTX-M-3 | 4 (2.9%) | 2 (2.5%) | 1.000 |
| CTX-M-32 | - | 2 (2.5%) | 0.136 |
| CTX-M-55 | 1 (0.7%) | 1 (1.2%) | 1.000 |
| CTX-M-61 | 1 (0.7%) | - | 1.000 |
| SHV-12 | 2 (1.4%) | - | 0.532 |
| SHV-2 | 1 (0.7%) | - | 1.000 |
| SHV-5 | 4 (2.9%) | 9 (11.1%) | 0.018* |
| SHV-7 | - | 1 (1.2%) | 0.370 |
| TEM-12 | 1 (0.7%) | - | 1.000 |
| TEM-52 | 3 (2.2%) | 1 (1.2%) | 1.000 |
| Unknown (not available for genotype analysis) | 16 (11.6%) | 9 (11.1%) | 1.000 |

**Underlying conditions**

| | Bacteremia (n = 138) | Sepsis with organ failure (n = 81) |
|-----------------|----------------------|----------------------------------|
| Heart disease | 16 (11.6%) | 23 (28.4%) | 0.003* |
| Vascular disease | 24 (17.4%) | 21 (25.9%) | 0.166* |
| Neurologic disease | 14 (10.1%) | 7 (8.6%) | 0.815 |
| Chronic pulmonary disease | 17 (12.3%) | 19 (23.5%) | 0.038* |
| Connective tissue disease | 3 (2.2%) | - | 0.298 |
| Ulcer disease | 6 (4.3%) | 2 (2.5%) | 0.713 |
| Liver disease | 16 (11.6%) | 25 (30.9%) | <0.001* |
| Diabetes mellitus | 34 (24.6%) | 20 (24.7%) | 1.000 |
| Moderate to severe renal disease | 44 (31.9%) | 62 (76.5%) | <0.001* |

(Continued)
Factors associated with in-hospital mortality (Cox-proportional hazard analysis)

The results of the univariate analysis on in-hospital mortality are displayed in Table 2. In order to assess the effect of the infecting organism (E. coli vs. K. pneumoniae) on mortality this parameter was also considered in the Cox-proportional hazard regression. The deceased had significantly higher CCIs. These patients also had significantly more often a hospital onset bacteremia, sepsis with organ failure, prior episodes of lower respiratory tract infection, intra-abdominal infection or primary bacteremia, underlying heart disease, liver- or renal disease. The survivors had more commonly a urinary tract infection prior to their bacteremia. The results of the Cox-proportional hazard regression (Table 4) showed that sepsis with organ failure was associated with a 4.5-fold higher hazard for mortality, renal disease and liver disease with 2.7-fold and 1.8-fold elevated hazard. The only protective factor was urinary tract infection that was associated with a hazard reduction for mortality of about 61% (HR 0.39). There was no statistically significant difference in the mortality risk between both species. Fig 2 shows the Kaplan Meier plot for cumulative survival stratified by ESBL-E species in relation to length of stay after onset of sepsis.

Discussion

In this study ESBL-KP-Bac bacteremia was associated with different origin, with sepsis with organ failure and younger age compared to ESBL-EC-Bac cases (S1 Table). Our results underline the findings of previous studies showing that K. pneumoniae infections are associated with more serious illness than E. coli infections [4, 6, 7, 22]. Even though K. pneumoniae bacteremia was not associated with delayed adequate anti-infective treatment it was associated with slightly increased mortality, but it did not reach the significance level. The small sub-cohort of ESBL-KP-Bac might explain this.

Several studies show that a delay of adequate antimicrobial chemotherapy is a risk factor for in-hospital death [1, 2, 4]. In our cohort, the association of sepsis with organ failure and reduced DAT demonstrate most likely the realization of the German sepsis guidelines [23]. Patients who present with severe sepsis or septic shock are recommended to be immediately initiated with an early initial broad-spectrum treatment including reserve antibiotics e.g. carbapenems. In our study, the presentation of severe sepsis happened before the initiation of the antimicrobial therapy. Hence, the observed association between disease severity and reduced DAT represents the response to the severity of the disease and not the cause of the disease. This refers to the observation that in sepsis with organ failure, an early effective antimicrobial therapy is associated with significantly reduced mortality risk [24]. Even though most cases of sepsis with organ failure received appropriate antimicrobial therapy within the first day, more than 25% showed a DAT of more than 1 day. This observation might at least partly explain the remaining high mortality (51.9%) in patients with severe sepsis and septic shock.
## Table 2. Univariate analysis of survivors and non-survivors after ESBL-E sepsis

| Parameter | Survived (n = 165) | Deceased (n = 54) | P-value |
|-----------|-------------------|------------------|---------|
| Age years / Age < 61 years | 80 (48.5%) | 29 (53.7%) | 0.534 |
| Male sex | 107 (64.8%) | 39 (72.2%) | 0.406 |
| Charlson comorbidity index > 6 | 5 (3; 8) | 8 (6; 10) | <0.001 |
| Days from admission to onset | 5 (1; 18) | 27 (15; 24) | <0.001 |
| Days from onset to discharge/death | 14 (9; 26) | 12 (2; 31) | 0.504 |
| Polymicrobial bacteremia | 23 (13.9%) | 6 (11.1%) | 0.817 |
| Hospital onset | 89 (53.9%) | 44 (81.5%) | <0.001* |
| ESBL colonization before onset | 106 (64.2%) | 32 (59.3%) | 0.520 |
| Bacteraemia due to *E. coli* | 122 (73.9%) | 38 (70.4%) | 0.601 |
| Bacteraemia due to *K. pneumoniae* | 43 (26.1%) | 16 (29.6%) | 0.647 |

### Origin of ESBL-E bacteraemia

| Parameter | Survived (n = 165) | Deceased (n = 54) | P-value |
|-----------|-------------------|------------------|---------|
| Urinary tract infection | 66 (40.0%) | 9 (16.7%) | 0.002* |
| Lower respiratory tract infection | 24 (14.5%) | 16 (29.6%) | 0.024* |
| Intra-abdominal infection | 11 (6.7%) | 11 (20.4%) | 0.007* |
| Surgical site infection | 4 (2.4%) | 4 (7.4%) | 0.105* |
| Primary bacteraemia | 10 (6.0%) | 20 (37.0%) | <0.001* |
| Other | 8 (4.8%) | - | 0.199 |
| Unknown Origin | 44 (26.7%) | 12 (22.2%) | 0.647 |

### Severity of illness and delay of anti-infective treatment

| Parameter | Survived (n = 165) | Deceased (n = 54) | P-value |
|-----------|-------------------|------------------|---------|
| Bacteremia/ sepsis | 126 (76.4%) | 12 (22.2%) | <0.001* |
| Severe sepsis/ septic shock | 39 (23.6%) | 42 (77.8%) | 0.028* |
| Delayed anti-infective treatment (days) | 1 (0;3) | 1 (0;2) | - |

### ESBL Genotype

| Parameter | Survived (n = 165) | Deceased (n = 54) | P-value |
|-----------|-------------------|------------------|---------|
| No ESBL genotype | 6 (3.6%) | 1 (1.9%) | 1.000 |
| CTX-M-1 | 31 (18.8%) | 8 (14.8%) | 0.682 |
| CTX-M-14 | 11 (6.7%) | 3 (5.6%) | 1.000 |
| CTX-M-15 | 71 (43.0%) | 27 (50.0%) | 0.431 |
| CTX-M-2 | 1 (0.6%) | - | 1.000 |
| CTX-M-2/97 | 2 (1.2%) | - | 1.000 |
| CTX-M-3 | 6 (3.6%) | - | 0.340 |
| CTX-M-32 | - | 2 (3.7%) | 0.060 |
| CTX-M-55 | 1 (0.6%) | 1 (1.9%) | 0.433 |
| CTX-M-61 | 1 (0.6%) | - | 1.000 |
| SHV-12 | 2 (1.2%) | - | 1.000 |
| SHV-2 | 1 (0.6%) | - | 1.000 |
| SHV-5 | 9 (5.5%) | 4 (7.4%) | 0.740 |
| SHV-7 | 1 (0.6%) | - | 1.000 |
| TEM-12 | 1 (0.6%) | - | 1.000 |
| TEM-52 | 3 (1.8%) | 1 (1.9%) | 1.000 |
| Unknown (not available for genotype analysis) | 18 (10.9%) | 7 (13.0%) | 0.631 |

### Underlying conditions

| Parameter | Survived (n = 165) | Deceased (n = 54) | P-value |
|-----------|-------------------|------------------|---------|
| Heart disease | 19 (11.5%) | 20 (37%) | >0.001* |
| Vascular disease | 30 (18.2%) | 15 (27.8%) | 0.173* |
| Neurologic disease | 18 (10.9%) | 3 (5.6%) | 0.299 |
| Chronic pulmonary disease | 24 (14.5%) | 12 (22.2%) | 0.206 |
| Connective tissue disease | 3 (1.8%) | - | 1.000 |

(Continued)
ESBL-KP-Bac cases were associated with sepsis with organ failure. However, they did not show significant differences in comorbidities compared to ESBL-EC-Bac cases (S1 Table). This might be explained by a potentially higher virulence of *K. pneumoniae* compared to *E. coli*. An earlier study on length of hospital stay included parts of the data at hand. Altogether 1,851 cases of bacteremia with (ESBL-positive and –negative) *Enterobacteriaceae* were analyzed then [10]. In that study, *K. pneumoniae* cases were associated with significantly prolonged hospital stay compared to *E. coli* cases. This most likely indicates a more problematic course of infection in *K. pneumoniae* cases. However, in that former study no data on antimicrobial therapy was analyzed [10]. Our present results support the previous findings after adjustment for timely and adequate antimicrobial therapy.

### Table 2. (Continued)

| Parameter                          | Survived (n = 165) | Deceased (n = 54) | P-value |
|------------------------------------|--------------------|-------------------|---------|
| Ulcer disease                      | 5 (3.0%)           | 3 (5.6%)          | 0.411   |
| Liver disease                      | 20 (12.1%)         | 21 (38.9%)        | >0.001* |
| Diabetes mellitus                  | 36 (21.8%)         | 18 (33.3%)        | 0.103*  |
| Moderate to severe renal disease   | 61 (37.0%)         | 45 (83.3%)        | >0.001* |
| Cancer/immunological disease       | 66 (40.0%)         | 21 (38.9%)        | 1.000   |

Continuous parameter are displayed as median (interquartile range), categorical parameter as number (percentage). ESBL, extended-spectrum beta-lactamase.

*, parameter was included in the Cox regression analysis on risk factors for death.

doi:10.1371/journal.pone.0158039.t002

### Table 3. Results of the multivariable binary logistic regression analysis on risk factors for sepsis with organ failure

| Parameter                          | P-value | OR | Upper—Lower CI 95 |
|------------------------------------|---------|----|-------------------|
| Moderate to severe renal disease   | <0.001  | 6.835 | 3.485–13.405     |
| Liver disease                      | 0.004   | 3.347 | 1.463–7.658      |
| Bacteremia due to *E. coli*        | <0.001  | 1 = reference |               |
| Bacteremia due to *K. pneumoniae*  | 4.499 | 2.168–9.337 |           |

OR, odds ratio. CI 95, 95% confidence interval.

doi:10.1371/journal.pone.0158039.t003

### Table 4. Results of the Cox-proportional hazard regression on clinical risk factors for death after ESBL-E sepsis.

| Parameter                          | P-value | HR | Upper—Lower CI 95 |
|------------------------------------|---------|----|-------------------|
| Bacteremia due to *E. coli*        | 0.060   | 1 = reference |               |
| Bacteremia due to *K. pneumoniae*  | 1.801   | 0.975–3.330 |           |
| Urinary tract infection            | 0.007   | 0.360 | 0.172–0.754      |
| Bacteremia/sepsis                  | <0.001  | 1 = reference |               |
| Severe sepsis/septic shock         | 4.543   | 2.134–9.673 |           |
| Liver disease                      | 0.045   | 1.801 | 1.013–3.203      |
| Moderate/ severe renal disease     | 0.016   | 2.675 | 1.198–5.973      |

HR, hazard ratio. CI 95, 95% confidence interval.

doi:10.1371/journal.pone.0158039.t004
The primary source of bacteremia differed significantly between both pathogens. While *E. coli* bacteremia was mostly found secondary to a urinary tract infection, *K. pneumoniae* cases were associated with surgical site infection (SSI), lower respiratory tract infection (LRTI) and unknown origin. Besides undetected colonization, the latter could be explained by health-care associated transmission. However, all sites of origin (SSI, LRTI and transmission) are likely since *K. pneumoniae* is often identified as outbreak pathogen [25, 26] and shows higher transmission potential than *E. coli* [27].

Urinary tract infection (UTI) as possible source of bacteremia was found less often associated with the development of a fatal bacteremia. This goes along with former studies on ESBL-E bacteremia [7, 28]. ESBL-E are common pathogens of urinary tract infections. A prior microbiologically diagnosed ESBL-E UTI might have supported an early initiation of anti-infective treatment at presentation of a secondary bacteremia.

Even though there are significant differences in the ESBL genotype distribution of both pathogens, none of these genotypes was associated with increased mortality. However, our data confirms that the most common genotypes among clinical ESBL-positive *E. coli* isolates in the United States and in Europe are currently CTX-M-15 and CTX-M-14 [29, 30]. In our *K.
pneumoniae isolates the most common ESBL genotypes were CTX-M-15 and SHV-5 which is also commonly observed in Europe [3, 31].

In our study, most of the isolates were reported resistant against piperacillin-tazobactam (pip-taz) due to their ESBL positivity. In 2011, CLSI recommended the interpretation of the breakpoint should be reported as found, irrespective of whether there was ESBL production [32]. Based on the current CLSI breakpoints, 35.2% of our isolates would be resistant to pip-taz. However, in this study, we focused on the results of the treatment based on the reported antibiogram. This goes along with the observed antimicrobial treatment showing carbapenems as mostly used agent, followed by quinolones.

This study has limitations. It was performed retrospectively and only ESBL-positive infections were assessed. Potential differences to ESBL-negative infections cannot be determined. The study was conducted at a single center. However, the data were collected from all patients within our hospital, regardless the respective department and likely represents the current course of these kind of infections in comparable hospitals. The species identification was performed using the Vitek 2 system. Since Vitek 2 cannot differentiate K.pneumoniae and K.variicola, it is possible that up to 10% of our analyzed K.pneumoniae isolates are K.variicola.

Future studies on carbapenem resistance should include ertapenem. Some isolates produce weak carbapenemases and may show decreased susceptibility to this substance only.

In conclusion, ESBL-positive K.pneumoniae bacteremia compared to ESBL-positive E.coli bacteremia is often associated with complicated infection and less often with uncomplicated infection such as urinary tract infection. In this small study group, pathogen on species level and genotypes were not associated with mortality, but with well-known factors as sepsis with organ failure. Knowledge of colonization and source of infection should be considered for empiric anti-infective treatment, especially in patients with septic shock to reduce DAT and mortality. Infections with ESBL-positive K.pneumoniae should be considered as more serious infections than comparable E.coli infections. Treatment and prevention strategies should be adjusted accordingly.

Supporting Information

S1 Table. Univariate analysis of clinical parameter in patients with ESBL-positive bacteremia due to E.coli in comparison to K.pneumoniae. Continuous parameter are displayed as median (interquartile range), categorical parameter as number (percentage). ESBL, extended-spectrum beta-lactamase.

(SDOCX)

S2 Table. Raw data.

(XLS)

Acknowledgments

We thank Sybille Müller-Bertling and Christoph Eller for excellent technical assistance. We furthermore thank Ivo Steinmetz for his laboratory support. This project was executed within the scope of the RESET research consortium and was funded partly by grants from the Federal Ministry of Education and Health, Germany (01KI1013H to RL and PG).

Author Contributions

Conceived and designed the experiments: CS SG YP RL. Performed the experiments: CS SG CK YP RL. Analyzed the data: RL FS MD PG CS SG. Contributed reagents/materials/analysis tools: CK YP AK. Wrote the paper: CS SG RL PG MD. Investigation: IS. Supervision: IS.
References

1. Rottier WC, Ammerlaan HS, Bonten MJ. Effects of confounders and intermediates on the association of bacteremia caused by extended-spectrum beta-lactamase-producing Enterobacteriaceae and patient outcome: a meta-analysis. The Journal of antimicrobial chemotherapy. 2012; 67(6):1311–20. Epub 2012/03/08. doi: 10.1093/jac/dks065 PMID: 22396430.

2. Schwaber MJ, Carmeli Y. Mortality and delay in effective therapy associated with extended-spectrum beta-lactamase production in Enterobacteriaceae bacteremia: a systematic review and meta-analysis. The Journal of antimicrobial chemotherapy. 2007; 60(5):913–20. Epub 2007/09/13. doi: 10.1093/jac/dkm318 PMID: 17848376.

3. Qureshi ZA, Paterson DL, Peleg AY, Adams-Haduch JM, Shutt KA, Pakstis DL, et al. Clinical characteristics of bacteremia caused by extended-spectrum beta-lactamase-producing Enterobacteriaceae in the era of CTX-M-type and KPC-type beta-lactamases. Clinical microbiology and infection: the official publication of the European Society of Clinical Microbiology and Infectious Diseases. 2012; 18(9):887–93. Epub 2011/09/29. doi: 10.1111/j.1469-0691.2011.03658.x PMID: 21951551; PubMed Central PMCID: PMC3252485.

4. Tumbarello M, Sanguinetti M, Montuori E, Trecarichi EM, Posteroaro B, Fiori B, et al. Predictors of mortality in patients with bloodstream infections caused by extended-spectrum-beta-lactamase-producing Enterobacteriaceae: importance of inadequate initial antimicrobial treatment. Antimicrobial agents and chemotherapy. 2007; 51(6):1987–94. Epub 2007/03/28. doi: 10.1128/aac.01509-06 PMID: 17387156; PubMed Central PMCID: PMC1891412.

5. Kumar A, Ellis P, Arabi Y, Roberts D, Light B, Parrillo JE, et al. Initiation of inappropriate antimicrobial therapy results in a fivefold reduction of survival in human septic shock. Chest. 2009; 136(5):1237–48. Epub 2009/08/22. doi: 10.1378/chest.09-0087 PMID: 19696123.

6. Chen SC, Wu WW, Yeh CH, Lai KC, Cheng KS, Jeng LB, et al. Comparison of Escherichia coli and Klebsiella pneumoniae liver abscesses. The American journal of the medical sciences. 2007; 334(2):97–105. Epub 2007/08/19. doi: 10.1097/MAJ.0b013e3181259c7 PMID: 17700198.

7. Ku NS, Kim YC, Kim MH, Song JE, Lee DH, Ahn JY, et al. Risk factors for 28-day mortality in elderly patients with extended-spectrum beta-lactamase (ESBL)-producing Escherichia coli and Klebsiella pneumoniae bacteremia. Archives of gerontology and geriatrics. 2014; 58(1):105–9. Epub 2013/08/31. doi: 10.1016/j.archger.2013.07.002 PMID: 23988261.

8. Kim SH, Kwon JC, Choi SM, Lee DG, Park SH, Choi JH, et al. Escherichia coli and Klebsiella pneumoniae bacteremia in patients with neutropenic fever: factors associated with extended-spectrum beta-lactamase production and its impact on outcome. Annals of hematology. 2013; 92(4):533–41. Epub 2012/11/20. doi: 10.1007/s00277-012-1631-y PMID: 23161391.

9. Peralta G, Lamelo M, Alvarez-Garcia P, Velasco M, Delgado A, Horcajada JP, et al. Impact of empirical treatment in extended-spectrum beta-lactamase-producing Escherichia coli and Klebsiella spp. bacteremia. A multicentric cohort study. BMC infectious diseases. 2012; 12:245. Epub 2012/10/09. doi: 10.1186/1471-2334-12-245 PMID: 23038999; PubMed Central PMCID: PMC3519701.

10. Leistner R, Gurntke S, Kohler C, Steinmetz I, Pfeifer Y, Eller C, Gastmeier P, et al. Molecular epidemiology of bloodstream infection. A multicentric cohort study. BMC infectious diseases. 2012; 12:245. Epub 2012/10/09. doi: 10.1186/1471-2334-12-245 PMID: 23038999; PubMed Central PMCID: PMC3519701.

11. Leistner R, Gurntke S, Kohler C, Steinmetz I, Pfeifer Y, Eller C, Gastmeier P, et al. Molecular epidemiology of bloodstream infection. A multicentric cohort study. BMC infectious diseases. 2012; 12:245. Epub 2012/10/09. doi: 10.1186/1471-2334-12-245 PMID: 23038999; PubMed Central PMCID: PMC3519701.

12. Leistner R, Gurntke S, Kohler C, Steinmetz I, Pfeifer Y, Eller C, Gastmeier P, et al. Molecular epidemiology of bloodstream infection. A multicentric cohort study. BMC infectious diseases. 2012; 12:245. Epub 2012/10/09. doi: 10.1186/1471-2334-12-245 PMID: 23038999; PubMed Central PMCID: PMC3519701.

13. Leistner R, Gurntke S, Kohler C, Steinmetz I, Pfeifer Y, Eller C, Gastmeier P, et al. Molecular epidemiology of bloodstream infection. A multicentric cohort study. BMC infectious diseases. 2012; 12:245. Epub 2012/10/09. doi: 10.1186/1471-2334-12-245 PMID: 23038999; PubMed Central PMCID: PMC3519701.

14. Leistner R, Gurntke S, Kohler C, Steinmetz I, Pfeifer Y, Eller C, Gastmeier P, et al. Molecular epidemiology of bloodstream infection. A multicentric cohort study. BMC infectious diseases. 2012; 12:245. Epub 2012/10/09. doi: 10.1186/1471-2334-12-245 PMID: 23038999; PubMed Central PMCID: PMC3519701.

15. Leistner R, Gurntke S, Kohler C, Steinmetz I, Pfeifer Y, Eller C, Gastmeier P, et al. Molecular epidemiology of bloodstream infection. A multicentric cohort study. BMC infectious diseases. 2012; 12:245. Epub 2012/10/09. doi: 10.1186/1471-2334-12-245 PMID: 23038999; PubMed Central PMCID: PMC3519701.
17. Hagel S, Brunhorst F. Sepsis [Article in German]. Intensivmed. 2011; 48:57–8. Epub 73. doi: 10.1007/s00390-010-0249-3
18. Grobner S, Linke D, Schutz W, Fladerer C, Madlung J, Autenrieth IB, et al. Emergence of carbapenem-nonsusceptible extended-spectrum beta-lactamase-producing Klebsiella pneumoniae isolates at the university hospital of Tubingen, Germany. Journal of medical microbiology. 2009; 58(Pt 7):912–22. Epub 2009/06/09. jmm.0.005850–0 [pii]doi: 10.1099/jmm.0.005850-0 PMID: 19502377.
19. Perez-Perez FJ, Hanson ND. Detection of plasmid-mediated AmpC beta-lactamase genes in clinical isolates by using multiplex PCR. Journal of clinical microbiology. 2002; 40(6):2153–62. Epub 2002/05/31. PMID: 12037080; PubMed Central PMCID: PMC130804.
20. Eller C, Simon S, Miller T, Frick JS, Prager R, Rabsch W, et al. Presence of beta-lactamases in extended-spectrum-cephalosporin-resistant Salmonella enterica of 30 different serovars in Germany 2005–11. The Journal of antimicrobial chemotherapy. 2013; 68(9):1978–81. Epub 2013/05/16. doi: 10.1093/jac/dkt163 PMID: 23674765.
21. Clermont O, Bonaconsi S, Bingen E. Rapid and simple determination of the Escherichia coli phylogenetic net. Applied and environmental microbiology. 2000; 66(10):4555–8. Epub 2000/09/30. PMID: 11010916; PubMed Central PMCID: PMC29342.
22. Devrim I, Guldfan G, Gunay I, Agin H, Guven B, Yilmazer MM, et al. Comparison of in vitro activity of ertapenem with other carbapenems against extended-spectrum-beta-lactamase-producing Escherichia coli and Klebsiella species isolated in a tertiary children's hospital. Expert opinion on pharmacotherapy. 2011; 12(6):845–9. Epub 2011/02/18. doi: 10.1517/14656566.2011.559460 PMID: 21323503.
23. Reinhart K. S2-Leitlinie: Diagnose und Therapie der Sepsis: Georg Thieme Verlag; 2007.
24. Kumar A, Roberts D, Wood KE, Light B, Parrillo JE, Sharma S, et al. Duration of hypotension before initiation of effective antimicrobial therapy is the critical determinant of survival in human septic shock. Critical care medicine. 2006; 34(6):1589–96. Epub 2006/04/21. doi: 10.1097/01.ccm.0000217961.75225.e9 PMID: 16625170.
25. Epstein EE, Pisney LM, Wendt JM, MacCannell DR, Janelle SJ, Kitchel B, et al. Carbapenem-resistant Klebsiella pneumoniae producing New Delhi metallo-beta-lactamase at an acute care hospital, Colorado, 2012. Infection control and hospital epidemiology: the official journal of the Society for Healthcare Epidemiologists of America. 2014; 35(4):390–7. Epub 2014/03/08. doi: 10.1086/675607 PMID: 24602944.
26. Lubbert C, Lippmann N, Busch T, Kaisers UX, Ducomble T, Eckmanns T, et al. Long-term carriage of Klebsiella pneumoniae carbapenemase-2-producing K pneumonia after a large single-center outbreak in Germany. American journal of infection control. 2014; 42(4):376–80. Epub 2014/04/01. doi: 10.1016/j.ajic.2013.12.001 PMID: 24679563.
27. Hilty M, Betsch BY, Bogli-Stuber K, Heiniger N, Stadler M, Kuffer M, et al. Transmission dynamics of extended-spectrum beta-lactamase-producing Enterobacteriaceae in the tertiary care hospital and the household setting. Clinical infectious diseases: an official publication of the Infectious Diseases Society of America. 2012; 55(7):967–75. Epub 2012/06/22. doi: 10.1093/cid/cis851 PMID: 22718774; PubMed Central PMCID: PMC3436924.
28. Tumbarello M, Sali M, Trecarichi EM, Leone F, Rossi M, Fiori B, et al. Bloodstream infections caused by extended-spectrum-beta-lactamase- producing Escherichia coli: risk factors for inadequate initial antimicrobial therapy. Antimicrobial agents and chemotherapy. 2008; 52(9):3244–52. Epub 2008/07/02. doi: 10.1128/aac.00063-08 PMID: 18591273; PubMed Central PMCID: PMC2533461.
29. Chen LF, Freeman JT, Nicholson B, Keiger A, Lancaster S, Joyce M, et al. Widespread dissemination of CTX-M-15 genotype extended-spectrum-beta-lactamase-producing enterobacteriaceae among patients presenting to community hospitals in the southeastern United States. Antimicrobial agents and chemotherapy. 2014; 58(2):1200–2. Epub 2013/11/20. doi: 10.1128/aac.01099-13 PMID: 24247126; PubMed Central PMCID: PMC3910860.
30. Brolund A, Edquist PJ, Makitalo B, Olsson-Liljequist B, Soderblom T, Wisell KT, et al. Epidemiology of extended-spectrum beta-lactamase-producing Escherichia coli in Sweden 2007–2011. Clinical microbiology and infection: the official publication of the European Society of Clinical Microbiology and Infectious Diseases. 2014; 20(6):O344–52. Epub 2013/10/15. doi: 10.1111/1469-0691.12413 PMID: 24118431.
31. Tumbarello M, Spanu T, Sanguinetti M, Citton R, Montuori E, Leone F, et al. Bloodstream infections caused by extended-spectrum-beta-lactamase-producing Klebsiella pneumoniae: risk factors,
molecular epidemiology, and clinical outcome. Antimicrobial agents and chemotherapy. 2006; 50 (2):498–504. Epub 2006/01/27. doi: 10.1128/aac.50.2.498-504.2006 PMID: 16436702; PubMed Central PMCID: PMCPmc1366869.

32. CLSI. Performance Standards for Antimicrobial Susceptibility Testing; 20th Informational Supplement (June 2010 Update). CLSI document M100-S20-U. 2010.