BMJ Open New methodology to assess the excess burden of antibiotic resistance using country-specific parameters: a case study regarding E. coli urinary tract infections

Noortje Grejanne Godijk,¹ Scott A McDonald ¹,² Wieke Altorf-van der Kuil,² Annelot F Schoffelen,² Eelco Franz,² Martin C J Bootsma^{1,3}

ABSTRACT

Objectives Antimicrobial resistant (AMR) infections are a major public health problem and the burden on population level is not yet clear. We developed a method to calculate the excess burden of resistance which uses countryspecific parameter estimates and surveillance data to compare the mortality and morbidity due to resistant infection against a counterfactual (the expected burden if infection was antimicrobial susceptible). We illustrate this approach by estimating the excess burden for AMR (defined as having tested positive for extended-spectrum beta-lactamases) urinary tract infections (UTIs) caused by E. coli in the Netherlands in 2018, which has a relatively low prevalence of AMR E. coli, and in Italy in 2016, which has a relatively high prevalence.

Design Excess burden was estimated using the incidence-based disability-adjusted life-years (DALYs) measure. Incidence of AMR E. coli UTI in the Netherlands was derived from ISIS-AR, a national surveillance system that includes tested healthcare and community isolates, and the incidence in Italy was estimated using data reported in the literature. A systematic literature review was conducted to find country-specific parameter estimates for disability duration, risks of progression to bacteraemia and mortality.

Results The annual excess burden of AMR E. coli UTI was estimated at 3.89 and 99.27 DALY/100 0000 population and 39 and 2786 excess deaths for the Netherlands and Italy, respectively,

Conclusions For the first time, we use country-specific and pathogen-specific parameters to estimate the excess burden of resistant infections. Given the large difference in excess burden due to resistance estimated for Italy and for the Netherlands, we emphasise the importance of using country-specific parameters describing the incidence and disease progression following AMR and susceptible infections that are pathogen specific, and unfortunately currently difficult to locate.

INTRODUCTION

Information on incidence and burden of disease (BoD) of infections with antimicrobialresistant (AMR) bacteria is valuable for setting public health priorities, designing and evaluating interventions.¹ However, such

STRENGTHS AND LIMITATIONS OF THIS STUDY

- \Rightarrow The strength of this study method is the application of the novel method to estimate the excess burden of an infection in two example countries to demonstrate its use.
- \Rightarrow We used country-specific and pathogen-specific parameters to estimate the excess burden of disease (BoD).
- \Rightarrow National-level surveillance data of the Netherlands informed the estimation of the incidence of resistant E. coli urinary tract infections.
- \Rightarrow The main limitation was that assumptions had to be made for some country-specific parameters for which no suitable studies were found: this might have affected the estimated difference in the burden and excess burden between the Netherlands and Italy.
- \Rightarrow Most parameter estimates used in the calculation of excess BoD were derived from studies in hospital populations whereas data from studies in the general population could lead to more accurate and better generalisable estimates.

mining, AI training, information is scarce,² even though AMR has been identified in the European Union/ European Economic Area (EEA) as a major public health problem.³

To gain insight into the AMR-associated BoD, composite health measures, such as the disability-adjusted life-years (DALYs) measure, which can be derived from clinical pathway progression models, and suitable data on mortality and morbidity⁴⁵ are useful. Composite health measures allow diseases and their infectious causes to be ranked in terms of burden,⁶ and—particularly if based on incidence data-also facilitate measurement of the impact of public health interventions. In the case of AMR, the DALY approach can also be applied to compare the burden across resistant infectious agents, between countries or regions, and across time.

to text

and

data

1

Protected by copyright, including for uses related

McDonald SA. Altorf-van der Kuil W, et al. New methodology to assess the excess burden of antibiotic resistance using countryspecific parameters: a case study regarding E. coli urinary tract infections. BMJ Open 2023;13:e064335. doi:10.1136/ bmjopen-2022-064335

To cite: Godijk NG,

Prepublication history and additional supplemental material for this paper are available online. To view these files, please visit the journal online (http://dx.doi.org/10.1136/ bmjopen-2022-064335).

Received 03 June 2022 Accepted 03 February 2023

Check for updates

C Author(s) (or their employer(s)) 2023. Re-use permitted under CC BY-NC. No commercial re-use. See rights and permissions. Published by BMJ.

¹Julius Center for Health Sciences and Primary Care, University Medical Centre Utrecht, Utrecht, The Netherlands ²Centre for Infectious Disease Control, National Institute for Public Health & the Environment, Bilthoven. The Netherlands ³Department of Mathematics, Utrecht University, Utrecht, The Netherlands

Correspondence to

Dr Noortje Grejanne Godijk; n.g.godijk@gmail.com



Figure 1 Outcome trees(s) for urinary tract infection (UTI), for antimicrobial-susceptible (upper panel) and antimicrobial-resistant (lower panel) infection. Transition probabilities (P) stratified by type of infection ([S]usceptible or [R]esistant) are indicated for several transitions, as are disability durations (DD).

Attempts to comprehensively estimate the BoD of resistant infection using DALY have only recently been published, and report a large burden of resistance.² To calculate BoD, parameters for, among others, the chance of progression from acute infection to severe health outcomes, the risk of mortality and duration in each health outcome are needed. These parameter values are needed for AMR and antimicrobial susceptible (AMS) infections separetely because some previous studies observed worse outcomes for AMR infections. On the other hand, a study on complicated P. aeruginosa urinary tract infections (UTIs) and multidrug resistance did not find a difference in 30-day mortality and another study on bacteraemic UTI did also not find an association between 30-day mortality and resistant profiles.^{3 7} Parameters to calculate the BoD using the DALY measures should be chosen based on study findings of specific pathogens and infection site to provide more insight on whether resistance increases BoD. Moreover, estimating the BoD brings conceptual challenges, such as determining to what health state resistant infections should be compared, as discussed previously by de Kraker and Lipsitch. For instance, AMR infections can be compared with AMS infections or to the situation in which the infections do not occur and the choice of comparison method influences the calculated excess harm caused by resistance.⁸

The aim of this paper is to introduce a method to calculate the *excess* BoD. By 'excess BoD' we mean the mortality and morbidity (computed as DALY) associated with resistance, over and above the mortality and

morbidity associated with infection by the same-but AMS pathogen. In this approach, AMS infections with incidence identical to that for AMR infections serve as a counterfactual to estimate the additional health burden that is attributable to resistance. Our approach is new in that we combine country-specific incidence numbers from surveillance data with country-specific parameter values to calculate the excess BoD for infection caused by a specific resistant pathogen. Methods in previous studies did not include country-specific and pathogen-specific data to estimate the BoD. Subsequently, the method is demonstrated by calculating the excess BoD for a single infection site (UTI) and a single bacterial agent (E. coli) as AMR compared with AMS E. coli, where a resistant E. coli UTI is defined as a tested urine sample containing E. coli which produce extended spectrum beta-lactamases (ESBLs) as confirmed by a laboratory. The excess BoD of these infections was assessed for two countries: Italy, which was previously estimated to have the highest antibiotic-resistant BoD in the EEA, and the Netherlands, which **g** was ranked third from last in the list of highest antibiotic resistant BoD in the EEA.² Note that our goal is to illustrate how the methodology can be applied to countries with differing AMR E. coli prevalence and with differing surveillance data available, and not to conduct a formal comparison of these countries in terms of excess burden. We selected UTIs because they are among the most frequent infections in both the outpatient and inpatient setting and we choose E. coli UTIs specifically because UTIs are frequently caused by E. coli.^{9 10} Furthermore, UTI is a

	Netherlands		Italy				
Parameter	Susceptible	Resistant	Susceptible	Resistant			
P(Bact UTI)	3.6% (95% CI 3.4% to 3.8%) ³⁹ *	3.6% (95% CI 3.4% to 3.8%) ³⁹ *	3.6% (95% Cl 3.4% to 3.8%)*	3.6% (95% CI 3.4% to 3.8%)*			
P(Death Bact)	11.3% (24/212) ²⁹	27.5% (19/69) ²⁹	5.47%†	26.2% ⁴⁰			
P(PTSD Bact)	Uniform(0.13, 0.21) ¹⁴	Uniform(0.13, 0.21) ¹⁴	Uniform(0.13, 0.21) ¹⁴	Uniform(0.13, 0.21) ¹⁴			
P(CogImp Bact)	Uniform(0.11–0.47) ¹⁴	Uniform(0.11–0.47) ¹⁴	Uniform(0.11–0.47) ¹⁴	Uniform(0.11–0.47) ¹⁴			
P(PhysImp Bact)	1.0 ¹⁴	1.0 ¹⁴	1.0 ¹⁴	1.0 ¹⁴			
P(Renal Bact)	Uniform(0.009–0.13) ¹⁴	Uniform(0.009–0.13) ¹⁴	Uniform(0.009–0.13) ¹⁴	Uniform(0.009–0.13) ¹⁴			
DD(UTI)	5.1d (95% CI 4.3 to 5.9) ⁴¹	8.7d (95%Cl 7.0 to 10.8) ⁴¹	10d (IQR (7–17)) ^{42 43}	10d (IQR (7–17) ^{42 43}			
DD(Bact)	2.9d (95% Cl 1.7 to 4.0) ⁴⁴	7.9d (95% CI 3.5 to 13.0) ⁴⁴	13±9 ⁴⁵	20±17 days ⁴⁵			
DW(UTI)	Uniform(0.039, 0.152) ¹⁴	Uniform(0.039, 0.152) ¹⁴	Uniform(0.039, 0.152) ¹⁴	Uniform(0.039, 0.152) ¹⁴			
DW(Bact)	Pert(0.579,0.655,0.727) ¹⁴	Pert(0.579,0.655,0.727) ¹⁴	Pert(0.579,0.655,0.727)14	Pert(0.579,0.655,0.727)14			
DW(PTSD)	Pert(0.07,0.808,0.108) ¹⁴	Pert(0.07,0.808,0.108) ¹⁴	Pert(0.07,0.808,0.108) ¹⁴	Pert(0.07,0.808,0.108) ¹⁴			
DW(CogImp)	Pert(0.026,0.043,0.064) ¹⁴	Pert(0.026,0.043,0.064)14	Pert(0.026,0.043,0.064)14	Pert(0.026,0.043,0.064)14			
DW(PhysImp)	Uniform(0.011,0.053) ¹⁴	Uniform(0.011,0.053) ¹⁴	Uniform(0.011,0.053) ¹⁴	Uniform(0.011,0.053) ¹⁴			
DW(Renal)	Uniform(0.03,0.487) ¹⁴	Uniform(0.03,0.487) ¹⁴	Uniform(0.03,0.487) ¹⁴	Uniform(0.03,0.487) ¹⁴			

Disease burden model parameter values, with references, for susceptible and resistant E. coli urinary tract infections Table 1 in the Netherlands and the Italy settings, as derived from systematic review

*Pooled value from Mangen et al.5

†Calculated using the mortality rate of resistant E. coli bacteraemia given in Palacios-Baena et al⁴⁰ and the ratio between resistant E. coli bacteraemia mortality and E.coli bacteraemia mortality in Tumbarello et al.45

CL confidence interval.

common cause of sepsis a life-threatening complication with a very high mortality rate for all ages.¹¹ The excess BoD for AMR E. coli has not been estimated previously for the Netherlands and Italy using national-level data and country-specific parameter values.

METHODS

We begin by reviewing the parameter requirements for DALY estimation, then describe the systematic reviews that were carried out to locate country-specific parameter values, and finally detail the calculation of AMR E. coli UTI incidence for both target countries.

Outcome trees

We modified an existing outcome tree (OT) developed by the European Centre of Disease Control (ECDC) describing the clinical progression pathway for UTI,² shown in figure 1. We describe the separate transition probability parameters, disability durations (DDs), and disability weights (DWs) that are needed to quantify the BoD, in DALYs, due to infection with either the susceptible or resistant strain as shown in figure 1. The method simulates an incidence of AMS E. coli that is equal to resistant E. coli to estimate what the additional burden would be of resistant E.coli compared with the same number of AMS E. coli infections. Our excess BoD approach involves subtracting the estimated annual DALY for AMS UTIs, using the 'susceptible' version of the OT, from the annual DALY for AMR E. coli UTIs, using the 'resistant' version

of the OT, while simulating that incidence is identical. We simulate this identical incidence for calculating the excess burden, because we assume that a person would have had a susceptible infection in case they would not have had a resistant infection. Thus, only the OT parameters for resistant and susceptible E. coli UTIs differ.

The starting health outcome of the OT is a symptomatic UTI, after which patients can recover, or progress to ⊳ secondary bacteraemia, and following bacteraemia progtraining, and ress to several long-term sequelae or death.

DALY parameters and calculation

The principal 'input' to the DALY computation is the number of incident cases, in the current example the number of people experiencing an AMR E. coli UTI in 1 year. Transition probabilities between symptomatic UTI and all subsequent health outcomes are required. These estimates are required for AMR and AMS *E. coli* UTI sepa-rately because the probability of transitioning from one **Q** health state to another is often not the same for AMR and AMS infections. We use the notation P(*Outcome* | *Outcome*) to indicate the progression probability from *Outcome*₁ to Outcome_o. For instance, P(BactlUTI) is the probability of progression to bacteraemia given symptomatic UTI. No mortality risk is assumed following a UTI that does not progress to secondary bacteraemia. The OT specifies mortality risk as the parameter P(DeathlBact).

In general, DALYs are calculated as follows: the years of life lost (YLL) are added to the total years lost due to

and

Protected by copyright, including for uses related to tex



Figure 2 YLD and YLL due to resistant and counterfactual susceptible E. coli urinary tract infections in the Netherlands in 2018. Lines indicate 95% uncertainty intervals. DALY, disability-adjusted life-year; YLD, years lost due to disability; YLL, years of life lost.

disability (YLD) which is calculated by summing over the YLD for each (non-fatal) health outcome in the OT: DAIV = VII + VID

$$\text{YLD}_i = \sum_i N_i \times \text{DW}_i \times \text{DD}_i$$

YLL=No. deaths×life expectancy (LE) at age of death. *N*=the yearly incidence of health outcome *i*.

DW = the average disability weight of health outcome *i*. DD=Average duration of disability *i*.

DALY combines the YLL due to premature mortality and YLD, which captures time lived by an individual in less than full health. A loss of 1 year of full health is equivalent to one DALY.¹² For the computation of YLDs, DWs and DDs for each health outcome are required. Given availability of hospital length of stay (LOS) data in the literature, LOS data can serve as a measure of DD if the health state can involve hospital stay. When a patient can transition to more than one, simultaneously experienced,

health outcome (so-called 'internal comorbidity'), such as the long-term sequelae following secondary bacteraemia (figure 1), DWs of the overlapping health outcomes can be adjusted to take this into account.¹³ We decided a priori to adopt the same DWs as used by ECDC.²¹⁴

The risk of recurrent UTI episodes per patient was incorporated using a simple multiplier approach. Dealing with recurrence is necessary as the incidence data consist of the number of patients with at least one UTI episode in 1 year, and the transition probability from UTI to otected bacteraemia is defined per patient, but the annual BoD will depend on the total number of episodes in a year. Therefore, given an average annual number of episodes Š per patient, *j*, the total duration of time spent in the health outcome symptomatic UTI in a year is defined as 8 *i*×DD[UTI].

yright, For the computation of YLL, normative LE values by age-group at death are needed. Consistent with previous BoD exercises,^{2 15} we chose to use the Global Burden of Disease project (GBD-2010)¹⁶ values.

All BoD measures were estimated using pre-existing software, the BCoDE toolkit V.1.4.¹⁷ In this software, Monte-Carlo simulation with 1000 iterations is employed uses related to compute 95% uncertainty intervals around the BoD. We present the excess BoD and resistant BoD as DALY per 100000 population (to allow comparison between countries), DALY per 100 cases (for assessing the patientlevel burden; also useful for between-country comparð ison), YLDs and YLL. text and

Systematic reviews

We performed systematic literature reviews to locate parameter estimates for the risk of progression to bacteraemia, risk of progression to health states following bacteraemia, LOS, other indicators of DDs and mortality



Figure 3 DALYs attributable to six sequelae of resistant and counterfactual susceptible E. coli urinary tract infections in the Netherlands in 2018. DALY, disability-adjusted life-year; PTSD, post-traumatic stress disorder.





Figure 4 Disability-adjusted life-years of resistant and counterfactual susceptible E. coli urinary tract infections in the Netherlands in 2018 per age- and sex-stratified group.

risk. The systematic reviews, performed separately for the Netherlands and Italy, are described in detail in online supplemental appendices 1-3, figures S1 and S2.

AMR E. coli UTI incidence in the Netherlands

Data of 2018 from ISIS-AR, a laboratory based AMR surveillance system in the Netherlands¹⁸ were used to estimate AMR E. coli UTI incidence. ISIS-AR contains results of antimicrobial susceptibility testing of bacterial isolates routinely tested in medical microbiology laboratories in the Netherlands. ISIS-AR contains data on all consecutive samples of patients, sampled in hospitals (inpatient and outpatient), general practices and long-term care facilities.¹⁹ The coverage of the surveillance system is shown in online supplemental figure S3. ISIS-AR contains data of 46 laboratory which represent around 80% of the Dutch hospitals.²⁰



Figure 5 YLD and YLL due to resistant and counterfactual susceptible E. coli urinary tract infections in Italy in 2016. Lines indicate 95% uncertainty intervals. DALY, disabilityadjusted life-year; YLD, years lost due to disability; YLL, years of life lost.

AMR E. coli UTI incidence was defined as the number of persons having at least one urinary AMR E. coli isolate in 2018 per 1000 population. The incidence was stratified by sex and 5-year age-group. Online supplemental table õ S1) shows the data used per sex and age-group to calculate the incidence and recurrence rate. Incidence is thus calculated as the total number of resistant E. coli UTI in 2018 per sex and age group divided by the number of inhabitants of the Netherlands per sex and age group in $\mathbf{\bar{s}}$ 2018, and subsequently multiplied by 1000. An algorithm was created which calculated the days in between two urinary test samples of the same patient to determine if two consecutive tests had been conducted within 2weeks in the same patient. If the urinary samples were more than 2weeks apart, the UTI was labelled as recurrent and then õ only one isolate was counted. If two tests conducted for the same individual were more than 2weeks apart, the UTI was defined as recurrent. As a sensitivity analyses, we also show the incidence if we would have defined a recurrent UTI as being longer than 3 months apart. We estimated the average number of recurrent episodes per patient per technologies year. Moreover, we estimate the total incidence of E. coli UTIs regardless of resistance to indicate the percentage of resistant E. coli UTIs in 2018. The analysis to estimate the incidence were performed in R V.4.0.2.

Estimation of AMR E. coli UTI incidence in Italy

No Italian source comparable to ISIS-AR was found. Therefore, we took seven steps to calculate the incidence.

Step 1. We took the number of UTIs (n=57271)reported in a study that retrospectively used primary care electronic medical records of around 1.1 million Italian general practitioner (GP) patients from 1 January 2016 through 31 December 2016.²¹ The coverage of this study

e



Figure 6 DALYs attributable to six sequelae of resistant and counterfactual susceptible E. coli urinary tract infections in Italy in 2016. DALY, disability-adjusted life-year; PTSD, post-traumatic stress disorder.

around $2\%^{22}$ and the Italian population size in 2016 reported on ISTAT was used to estimate the total number of patients with a UTI in the entire population in 2016.²²

Step 2. The sex and age-group distribution from a study on UTIs in 2015–2019 in an academic Italian high-volume centre, namely the University Hospital 'San Giovann di Dio e Ruggi d'Aragona' in Salerno, was used to distribute the total estimated UTIs among women (62.33%), men (37.77%) and age-groups.²³

Step 3. The number of E. coli UTIs was calculated assuming that 59.9% of UTIs were caused by E. coli as reported in Cardone *et al*²⁴ which we identified in the systematic review (online supplemental appendix 1).²⁴ From January 2013 to June 2017, Cardone *et al*²⁴ included urine samples collected in the emergency department and used two inclusion criteria. The urine samples had to be collected in (1) patients with UTI symptoms and (2) it had to be their first positive culture urine culture in a given year.

Step 4. A large study from April 2007 to April 2008 in 20 microbiology laboratories found that 15.1% of E. coli bacteraemia produced ESBL²⁵ and this percentage was



Figure 7 Disability-adjusted life-years of resistant and counterfactual susceptible E. coli urinary tract infections in Italy in 2016 per age- and sex-stratified group.

ated

to text

t and

data mining, AI training, and similar technologies

 Table 2
 Sex-aggregated and age-aggregated YLD, YLL and DALY estimates for antimicrobial resistant and the counterfactual susceptible *E. coli* urinary tract infection, and estimated excess burden attributable to resistance (in DALY), for the Netherlands in 2018

	YLD (95% UI)	YLL (95% UI)	DALY (95% UI)	DALY/100 cases (95% UI)	DALY/100 000 pop (95% UI)
Resistant	458	1223	1581	20.84	9.20
	(424 to 497)	(1016 to 1234)	(1467 to 1701)	(19.34 to 22.42)	(8.58 to 9.90)
Counterfactual susceptible	445	467	913	12.03	5.31
	(409 to 482)	(424 to 513)	(854 to 934)	(11.26 to 12.84)	(4.97 to 5.67)
Excess burden	13	655	669	8.81	3.89

DALY, disability-adjusted life-year; pop, population; UI, uncertainty interval; YLD, years lost due to disability; YLL, years of life lost.

then applied to the results of Step 3 to estimate the AMR *E. coli* UTI incidence.

Step 5. To estimate the incident number of AMR *E. coli* UTIs per 5-year age category as needed for the BCoDE toolkit V.1.4¹⁷ (eg, 10–14, 15–19), we distributed UTIs within the age-categories used in Serretiello *et al*²³ proportionally according to the age-category and sex-specific population size.

Step 6. To calculate the incident number of AMR *E. coli* UTIs including clinical and outpatient cases, we used the same ratio of hospital to GP cases and outpatient to GP cases, sex and age-stratified, as in the Netherlands. We

used the same recurrence rate as we found in the Netherlands, as we were unable to identify a better estimate.

All calculations for the Italian incidence can be found online (https://github.com/NoorGo/ExcessBurden).

Patient and public involvement

There was no direct patient or public involvement in the design of this study.

 Table 3
 Incidence of resistant E. coli urinary tract infection including recurrent UTI in 2018 in the Netherlands and 2016 in Italy of females stratified by age

	Netherlands				Italy					
Age and sex category	Population (N)	Number of infections	Incidence rate	Incidence/ 100000 inhabitants	Population (N)	Number of infections	Incidence rate	Incidence/ 100000 inhabitants		
Females										
0	82565	10	0.00012	12.1	232955	6185	0.02655	2655.2		
1–4	340514	110	0.00032	32.3	1017487	8155	0.00801	801.5		
5–9	452 563	130	0.00029	28.7	1 385 255	1544	0.00111	111.5		
10–14	471 948	58	0.00012	12.3	1 384 866	1159	0.00084	83.7		
15–19	511 180	54	0.00011	10.6	1 391 122	2626	0.00189	188.8		
20–24	525964	121	0.00023	23.0	1 472 791	6411	0.00435	435.3		
25–29	545838	155	0.00028	28.4	1 607 399	6619	0.00412	411.8		
30–34	522235	131	0.00025	25.1	1 761 403	7940	0.00451	450.8		
35–39	512431	105	0.00020	20.5	2 037 299	10088	0.00495	495.2		
40–44	521 589	100	0.00019	19.2	2399975	13999	0.00583	583.3		
45–49	634635	173	0.00027	27.3	2490023	14392	0.00578	578.0		
50–54	635 623	227	0.00036	35.7	2420239	24738	0.01022	1022.1		
55–59	605380	362	0.00060	59.8	2110923	25965	0.01230	1230.0		
60–64	542 198	364	0.00067	67.1	1 891 237	26513	0.01402	1401.9		
65–69	503 662	388	0.00077	77.0	1 927 499	29486	0.01530	1529.8		
70–74	447 439	499	0.00112	111.5	1 533 451	22993	0.01499	1499.4		
75–79	314838	540	0.00172	171.5	1 552 174	24926	0.01606	1605.9		
80–84	235 430	525	0.00223	223.0	1 227 709	18861	0.01536	1536.2		
85+	248011	820	0.00331	330.6	1 365 423	21841	0.01600	1599.6		

Table 4 Incidence of resistant E. coli urinary tract infection (UTI) including recurrent UTI in 2018 in the Netherlands and 2016 in Italy in males stratified per age

	Netherlands				Italy					
Age and sex category	Population (N)	Number of infections	Incidence rate	Incidence/ 100000 inhabitants	Population (N)	Number of infections	Incidence rate	Incidence/ 100 000 inhabitants		
Males										
0	87001	12	0.00014	13.8	246656	10516	0.04263	4263.2		
1–4	358019	21	0.00006	5.9	1075850	12419	0.01154	1154.3		
5–9	475503	10	0.00002	2.1	1469465	4714	0.00321	320.8		
10–14	494511	8	0.00002	1.6	1 469 325	850	0.00058	57.8		
15–19	536852	15	0.00003	2.8	1 490 426	1712	0.00115	114.9		
20–24	542817	15	0.00003	2.8	1 563 396	4037	0.00258	258.2		
25–29	560319	31	0.00006	5.5	1653304	3049	0.00184	184.4		
30–34	530 554	35	0.00007	6.6	1776419	3479	0.00196	195.8		
35–39	512925	19	0.00004	3.7	2043171	9548	0.00467	467.3		
40–44	516723	35	0.00007	6.8	2380558	4098	0.00172	172.2		
45–49	634188	69	0.00011	10.9	2441662	10417	0.00427	426.6		
50–54	644223	114	0.00018	17.7	2337449	11304	0.00484	483.6		
55–59	606130	163	0.00027	26.9	1990139	10322	0.00519	518.6		
60–64	537 540	216	0.00040	40.2	1755003	30703	0.01749	1749.5		
65–69	495875	349	0.00070	70.4	1757419	37111	0.02112	2111.7		
70–74	424 486	440	0.00104	103.7	1 322 775	21 430	0.01620	1620.1		
75–79	273902	437	0.00160	159.5	1227379	17312	0.01411	1410.5		
80–84	172825	357	0.00207	206.6	826785	13985	0.01691	1691.5		
85+	122648	368	0.00300	300.0	629140	8887	0.01413	1412.6		

RESULTS

The results of the systematic review are discussed in online supplemental appendix 4, and the identified parameter values are described in table 1.

Parameters

The Netherlands

P(DeathlBact) for AMS E. coli was 11.3% and for AMR E. coli 27.5%. We estimated the DD(UTI) for AMS E. coli at 5.1 days (95% CI 4.3 to 5.9) and for AMR E. coli at 8.7 days (95% CI 7.0 to 10.8). DD(Bact) for AMS E. coli is 2.9 days $(95\%\ {\rm CI}\ 1.7\ {\rm to}\ 4)$ and for AMR E. coli $7.9\ {\rm days}\ (95\%\ {\rm CI}\ 3.5$ to 13.0). All parameters and their sources can be found in table 1.

Italy

P(Death|Bact) for AMS E. coli was 5.47% and for AMR E. coli this was estimated to be 26.5%.⁵ We were only able to find a single Italian parameter value for DD(UTI), which did not distinguish between AMS E. coli and AMR E. coli (10.7 days, IQR (7-17)). DD(Bact) for AMS E. coli was estimated at 13 days (SD=9) and for AMR E. coli at 20 days (SD=17).

Excess burden

The Netherlands

data minin Per 100 0000 inhabitants we found an excess burden of 3.9 DALY/100 000. The YLL component accounted for ⊳ 98% of the excess BoD. We found 39 (59%) excess deaths compared with the AMS model. Figure 2 shows the YLL and YLD for the Netherlands, while assuming equal incidence of susceptible and AMR E. coli. Per 100 cases the excess burden was estimated at 8.8 DALY/100 cases. The greatest excess burden was observed for bacteraemia (658 DALY) as can be seen in figure 3 which shows the excess burden for each of the six specified health outcomes in the clinical pathway progression model for UTI. Sexgroup and age-group differences in both BoD and excess burden were apparent (figure 4); the latter was two times greater for females (527 compared with 257 DALY per year in the population of males).

Italy

Per 100000 inhabitants In Italy, we estimated an excess burden of 99 DALY/100 000. The YLL component accounted for 99.7% of the excess burden and 2786 (77.0%) excess deaths were estimated. Per 100 cases the excess BoD was estimated at 12.3 DALY/100 cases. Figure 5 shows the YLL and YLD for Italy for AMR E. coli UTI and when simulating equal incidence of the counterfactual

AMS E. coli UTI. Figure 6 which shows the excess burden for each of the six specified health outcomes in the clinical pathway progression model for UTI. Sex-group and age-group differences in both BoD and excess burden were apparent (figure 7); the excess burden was 1.3 times greater for females (34036 compared with 26184 DALY). The 5-year age-group contributing the largest estimated excess BoD was 55 to 59-year-old women and 65 to 69-year-old men (5990 and 6041 DALY, respectively).

Resistant burden

The Netherlands

In the Netherlands a total of 9623 AMR E. coli UTIs occurred in 2018 based on the tested isolates in ISIS-AR, corresponding to an annual incidence of 0.56 AMR E. coli UTIs/1000 inhabitants. This incidence includes recurrent UTIs. These UTIs occurred in 7586 unique patients, resulting in an annual incidence of 0.44 AMR E. coli UTIs/1000 inhabitants, excluding recurrent UTIs. Online supplemental table S1 was used to calculate the AMR E. coli UTI incidence and recurrence rate per age and sex group. Of the unique AMR E. coli UTIs, 64.2% occurred in women and 62.3% in people aged 65 years or older. The total number of E. coli UTI in 2018 was 199441 and excluding recurrent UTI 165 258. The incidence including recurrent UTIs was 11.61/1000 inhabitants and 9.62/1000 inhabitants excluding recurrent E. coli UTI. The percentage resistant E. coli UTIs was 4.8% including recurrent UTIs and 4.6% excluding recurrent UTIs of the total number of E. coli UTIs in 2018. Online supplemental table S2 was used to calculate the E. coli UTI incidence and recurrence rate per age and sex group. In the sensitivity analysis in which we assumed a recurrent UTI to be more than 3 months apart we found an overall incidence of 0.47 AMR E. coli UTIs/1000 inhabitants and an incidence of 0.44 AMR E. coli UTI/1000 inhabitants excluding recurrent UTIs. Online supplemental table S3 shows the data of the incidence calculation for the sensitivity analysis.

Per 100000 inhabitants in the Netherlands, we estimated an AMR E. coli UTI incidence of 9.2 DALY/100 000 inhabitants (95% UI 8.5 to 9.9). The YLL component accounted for 71.0% of the resistant BoD and 66 deaths

were estimated. The sex-aggregated and age-aggregated BoD for AMR E. coli UTI in the Dutch population in 2018 was estimated at 1581 DALY (95% UI 1467 to 1701), or per 100 cases 20.8 DALY (95% UI 19.3 to 22.3) DALY (table 2). The resistant BoD for females was approximately two times that for males (1011 compared with 570 DALY) as shown in figure 4. Figure 3 shows the BoD for the specified health outcomes in the UTI clinical pathway progression model. The health outcome with the highest BoD for UTIs caused by AMR E. coli was bacteraemia Protected by (1127 DALY, 95% UI 1020 to 1238).

Italv

In Italy in 2016, we estimated 490332 AMR E. coli UTI copy and an incidence of 8.1 UTIs/1000 inhabitants excluding recurrent UTI. In women, 56% of infections occurred and 44% occurred in people aged ≥65 years. Incidences per age and sex group can be found in tables 3 and 4.

inclu In Italy, we estimated 192 DALY/100 000 (95% UI 181 to 203). The YLL component accounted for 66.9% of the resistant UTI BoD. For the AMR model 3617 (95% UI 3352 to 3884) deaths were estimated. The sex-aggregated g and age-aggregated BoD for resistant AMR E. coli UTI in uses rela the Italian population in 2016 was estimated at 166488 (95% UI 109744 to 123106) DALY, or 23.8 DALY per 100 cases (table 5). Just as for the Netherlands, the health outcome with the highest BoD for UTIs caused by AMR te E. coli was bacteraemia (78 686 DALY, 95% UI 72 736 to 84 493), which also caused the larger excess burden (69885) e DALY) as can be seen in figures 3 and 6). The resistant an BoD for females was approximately 1.3 times that for males (64878 compared with 51610 DALY). The 55 to data 59-year-old women (9688 DALY) and 65 to 69-year-old men contributed the most (9765 DALY). Bululu

DISCUSSION

training We developed a method for estimating the excess BoD due to antimicrobial resistance, and applied the method to AMR E. coli UTI for two countries using country-specific , and parameters and incidence data. Using country-specific parameters for BoD estimates is crucial, as outcome measures (eg, mortality) are not only influenced by

accounted for 71	0% of the resistant l	BoD and 66 deaths	measures (eg, mo	ortality) are not o	only influenced by	similar te					
Table 5 Sex-aggregated and age-aggregated YLD, YLL and DALY estimates for resistant and counterfactual susceptible <i>E.</i> coli urinary tract infection, and estimated excess burden attributable to resistance (in DALY), for Italy in 2016 6000000000000000000000000000000000000											
	YLD (95% UI)	YLL (95% UI)	DALY (95% UI)	DALY/ 100 cases (95% UI)	DALY/ 100 000 pop (95% UI)	ogies.					
Resistant	38499.48	77989	116488	23.76	192.02						
	(35 387 to 41 684)	(72 056 to 83 785)	(109 744 to 123 106)	(22.38 to 25.11)	(180.90 to 202.92)						
Counterfactual susceptible	38349	17920	56268	11.48	92.75						
	(35 212 to 41 359)	(15 134 to 21 105)	(52 069 to 60 696)	(10.62 to 12.43)	(85.83 to 100.49)						
Excess burden	151	60 0 69	60220	12.28	99.27						
DALY, disability-adju	DALY, disability-adjusted life-year: pop. population: UI, uncertainty interval: YLD, years lost due to disability: YLL, years of life lost.										

BMJ Open: first published as 10.1136/bmjopen-2022-064335 on 18 December 2023. Downloaded from http://bmjopen.bmj.com/ on June 13, 2025 at Agence Bibliographique de Iseignement (ABES

≥

resistance itself, but can also be influenced by inappropriate treatment,⁸ and BoD depends on the prevalence of comorbidities, as well as country-specific differences in hospital and prevention policies.²⁶ Previous large BoD studies such as Cassini *et al*² did not use country-specific parameter estimates,² whereas our results indicate that this is important. Two examples, among others that we found in our study, of why the use of country-specific parameters is important are that parameters such as the risk of death following bacteraemia and the disease duration of bacteraemia we found in the literature differed between Italy and the Netherlands. Subsequently these parameter differences between Italy and the Netherlands contribute to the differences in the excess burden between Italy and the Netherlands.

YLL accounted for most of the estimated AMR BoD in the Netherlands and in Italy (71% and 66.3%, respectively). A previous study on healthcare-associated (HA) infections, including bloodstream infections and UTI, based on data of Italy in 2016, also found that the majority of the BoD of AMR was attributable to YLL (79.7%).²⁷ Regarding the burden of AMR in DALYs per 100000 population, HA UTIs were estimated at 81.2 (69.0-94.4) DALYs/100 000 population. Both studies noted that UTIs were the second¹⁴ or most frequent²⁷ HA in terms of incidence. The difference in excess BoD and in the AMR disease burden between the Netherlands and Italy that we found might be partly due to differences in treatment and resistance testing policies. Since our literature search, a Dutch study in eight hospitals was published suggesting a different mortality when comparing highly resistant to non-highly resistant bacteraemia, namely an RR of 1.08 (95% CI 0.48 to 2.41).²⁸ This estimated mortality would imply that our estimates of the excess burden for NL may be over-estimated as the mortality risk difference of Rottier *et al*²⁸ is smaller than that of van Hout *et al*.²⁹ However, the CI of Rottier *et al*²⁸ is relatively large and of the bacteraemia that were included, only 52% (n=1001) had the urinary tract as source and 62% (n=1190) was caused by E. coli.

Previous incidence estimates of resistant E. coli UTI based on data from 2015 indicate a third generation cephalosporin resistant E. coli UTI incidence in Italy that is 7.3 times higher than in the Netherlands, and a carbapenem resistant E. coli UTI incidence that is 12.3 times higher.² In the current study, we estimated AMR E. coli incidence to be 18.3 times higher in Italy in 2016 than in the Netherlands in 2018. However, these previous estimates from Cassini et al^2 were derived using a different approach²; namely, the incidence of blood-stream infection served as primary data, which was then extrapolated to specific infection sites and to each EU/EAA country. Also, in contrast to the study of Cassini *et al*,² we use country-specific parameters which might be more suitable to indicate differences between countries in contributors to BoD. In a recent burden study DALYs attributable to and associated with bacterial AMR for 23 pathogens and 88 pathogen-drug combinations in 204 countries and territories in 2019 are

€ from http://bmjopen.bmj.com/ on June 13, 2025 at Agence Bibliographique de (ABES) . ≥ <u>0</u>

provided. The authors mention the difficulty of understanding the burden of AMR when data are sparse and mention that because of data sparsity, they assumed the relative risk of death was the same for every syndrome, location and age group.³⁰ We also found it difficult to be treated as a sparse and the same for every syndrome, location and age group.³⁰ We also found it difficult to be treated as a sparse and have argued that such data is important for accurate excess burden estimation at country level be cause country-specific parameters of for example be accurate excess burden estimation at country level be cause counterfactual in the BoD calculation depend on the type of intervention.⁸ The excess BoD method proposed in the current study defines the susceptible counterfactual to have identical incidence as resistant infection. This method could accordingly be useful for a new antibiotic use, and a combination of intervention, and the replacement scenario of a the resistant strain. For these estimations, the model parameters could for example be adjusted at the resistant strain. For these estimations, the model parameters could for example be adjusted at the resistant strain. For these estimations, the model parameters could for example be adjusted at ³¹ Under the assumption that the replacement scenario (mostly) occurring, the comparison group should be the same group of patients with infections caused by ASM). Astrength of this study is that we used national-level for the surveillance data of the Netherlands to calculate the model parameters of the Netherlands to calculate the model we to exite the indicated of the Metherlands to calculate the model parameters and the replacement scenario (mostly) occurring, the comparison group should be the same group of patients with infections caused by ASM).

surveillance data of the Netherlands to calculate the a incidence of resistant E. coli UTI. The use of these data enabled us to estimate the incidence of AMR E. coli as a basis for the BoD estimate. However, the use of these data harbour some limitations. First, the national coverage is less than 100%; therefore AMR E. coli UTI incidence uning, is underestimated. Also, in Italy the study on which we based our estimation of the proportion of resistant E. coli is dependent on samples being taken, which is also sensitive to testing practice and does not have a complete national coverage. However, the BoD experienced by these 'missed' patients is expected to be small because their UTI resolved on first line treatment and therefore, they experienced little BoD. Their chance of progressing to bacteraemia would be minimal. Our DALY estimate is mostly determined by those patients that develop bacteraemia, which has an accompanying high risk of mortality. Second, the surveillance date are routine data from medical microbiological laboratories. The ISIS-AR data only contains UTIs that have been sampled and tested for resistance. In general practices in the Netherlands, UTIs are often sampled only when infection is not eliminated after initial treatment. A part of the UTIs, therefore, may have been missed in our study. However, since we based our calculations on AMR infections only, we do not expect that this has largely influenced our estimates.

Another strength of this study is that we not only propose a new method to calculate the excess BoD, but that we also apply our method to two countries to demonstrate its use and explore the methods drawbacks. A drawback of this method, as mentioned previously,³³ is that it often is difficult to locate high-quality AMR surveillance data and country-specific AMR attributable mortality and morbidity parameters, as we experienced in the current study. Even though we performed a systematic review, we were not able to locate relevant studies and/or recent estimates for all parameters. In low-income and middle-income countries data scarcity is an even larger problem, which makes using countryspecific parameter estimates and incidence data as we advise for out method harder, even though the use of country-specific parameters is probably even more important when comparing developing to developed countries. Apart from the higher percentage of resistance in Italy, the difference in parameter estimates between Italy and the Netherlands explain the larger BoD and excess BoD for Italy. For the Netherlands, available studies showed a smaller difference in the bacteraemia mortality rate for AMR E. coli and AMS E. coli (27.5% vs 11.3%, respectively) than for Italy (26.2% vs)5.5%, respectively). Moreover, for the Netherlands DDs for the UTI and bacteraemia health outcomes were shorter. However, we had to make multiple assumptions of the model parameters, especially for Italy, as countryspecific data were not available for all estimates. These assumptions may also affect the estimated difference in the burden and excess burden between the Netherlands and Italy. For example, we used the same ratio of hospital to GP cases and outpatient to GP cases for Italy as for the Netherlands because we could not find specific data for Italy. However, in both the Netherlands and Italy antibiotics are not sold over the counter (in Italy there are some exceptions, eg, when the drug is necessary in order not to interrupt the treatment of a chronic disease³⁴); thus prescriptions are required,^{34 35} and it is most common in both countries to first visit the GP, get treatment if necessary, and thereafter get additional care if needed. For this reason, we choose to use the same ratio of hospital to GP cases and outpatient to GP cases, even though there are some antibiotic prescription and treatment differences between the two countries. Furthermore, the estimated mortality following bacteraemia as a consequence of UTI was estimated to be 11.3% for AMS E. coli and for AMR E. coli 27.5% in the Netherlands,²⁹ whereas a previous study in Finland, Sweden and Canada found a mortality rate of 9.2% of E. coli BSI with third-generation cephalosporin susceptibility and a mortality of 14.1% of E. coli BSIs with thirdgeneration cephalosporin resistance.³⁶ As we found few parameter estimates that were country-specific, we were unable to, for example, do a small meta-analysis, and get more valid estimates. Thus, our results should be interpreted with caution. The codes used to calculate the incidence in Italy, the excel in which the figures

were created and the excel sheets used to calculated the excess burden are available on the Github repository (https://github.com/NoorGo/ExcessBurden).³

Moreover, the assumed 15.1% resistance prevalence E. coli UTIs in Italy is likely to be an underestimate, as other data from 2017 suggested around 75% of the E. coli isolates in Italy to be resistant to at least one antibiotic group and around 45% to be resistant to three or more antibiotic groups,³⁸ however the 2017 prevalence was not specific for UTIs and we preferred to use UTI-specific **_** AMR *E. coli* estimates. Future research would benefit from using more recent country-specific surveillance data, when it becomes available, to more accurately estimate ŝ AMR E. coli incidence.

In addition, parameter estimates were limited by generative restricted analysis of confounders.²⁶ We did, however, stratify our results for age and sex. Moreover, we adjusted generative results for age and sex. the risk of mortality following bacteraemia for age. Future research could use parameter estimates derived from the general population. Most estimates used in this study were derived from studies in hospital populations. Parameter estimates based on studies in the general population could lead to more accurate estimates that are better generalisable to the Dutch and Italian populations. For example, hospital patients presenting with a UTI may more likely progress to bacteraemia, due to an already weakened immune system, than individuals who present with a UTI at the GP. As we were unable to **6** at locate parameter estimates in the general population, we **6** at **6** locate parameter estimates in the general population, we also recommend future research to focus on estimating these parameters. An example of such a study could be following GP patients who have a confirmed AMR or AMS *E. coli* UTI to estimate the probability of progression to bacteraemia and subsequent mortality.

To conclude, for the first time, we use country-specific and pathogen-specific parameters to estimate the excess burden of resistant infections. Given the large excess burden difference between AMR E. coli and AMS E. coli UTI, we emphasise the importance of using countryspecific parameters describing the incidence and disease progression following resistant and susceptible infections pu similar technologies that are pathogen-specific. Unfortunately, these parameters are currently difficult to locate.

Contributors NGG, SAM and MCJB conceptualised the study. NGG conducted the literature review and performed the data analyses with the help of SAM. NGG generated the figures and drafted the manuscript. WA-vdK and AFS had access to the ISIS-AR data and supplied the required data for the incidence calculations. WA-vdK created online supplemental figure S3. SAM, MCJB, WA-vdK, AFS and EF reviewed the manuscript and performed a critical revision of the manuscript text to clarify the methodology. NGG is guarantor and is responsible for the overall content.

Funding This study was supported by the research project RADAR (Risk Assessment and Disease burden of Antimicrobial Resistance) funded through the One Health European Joint Programme by the EU's Horizon-2020 Research and Innovation Programme (grant 773830).

Map disclaimer The inclusion of any map (including the depiction of any boundaries therein), or of any geographic or locational reference, does not imply the expression of any opinion whatsoever on the part of BMJ concerning the legal status of any country, territory, jurisdiction or area or of its authorities. Any such

expression remains solely that of the relevant source and is not endorsed by BMJ. Maps are provided without any warranty of any kind, either express or implied.

Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

Patient consent for publication Not applicable.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available in a public, open access repository. The code used to calculate the incidence in Italy, the spreadsheet in which the figures were created and the spreadsheets used to calculated the excess burden are available in the Github repository https://github.com/NoorGo/ExcessBurden.

Supplemental material This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iD

Scott A McDonald http://orcid.org/0000-0003-0788-6011

REFERENCES

- Wernli D, Jørgensen PS, Harbarth S, et al. Antimicrobial resistance: the complex challenge of measurement to inform policy and the public. PLoS Med 2017;14:e1002378.
- 2 Cassini A, Högberg LD, Plachouras D, et al. Attributable deaths and disability-adjusted life-years caused by infections with antibioticresistant bacteria in the EU and the European economic area in 2015: a population-level Modelling analysis. *Lancet Infect Dis* 2019;19:56–66.
- 3 Eliakim-Raz N, Babitch T, Shaw E. n.d. Risk factors for treatment failure and mortality among hospitalized patients with complicated urinary tract infection: A multicenter retrospective cohort study. *Clin Infect Dis*
- 4 Kretzschmar M, Mangen M-JJ, Pinheiro P, et al. n.d. New methodology for estimating the burden of infectious diseases in Europe. *PLoS Med*;9:e1001205.
- 5 Mangen M-JJ, Plass D, Havelaar AH, et al. The Pathogen- and incidence-based DALY approach: an appropriated methodology for estimating the burden of infectious diseases. PLoS One 2013;8:e79740.
- 6 GBD 2015 Disease and Injury Incidence and Prevalence Collaborators. Global, regional, and national incidence, prevalence, and years lived with disability for 310 diseases and injuries, 1990-2015: a systematic analysis for the global burden of disease study 2015. *Lancet* 2016;388:1545–602.
- 7 Horcajada JP, Shaw E, Padilla B, et al. Healthcare-associated, community-acquired and hospital-acquired Bacteraemic urinary tract infections in hospitalized patients: a prospective Multicentre cohort study in the era of antimicrobial resistance. *Clin Microbiol Infect* 2013;19:962–8.
- 8 de Kraker MEA, Lipsitch M. Burden of antimicrobial resistance: compared to what *Epidemiol Rev* 2021;43:53–64.
- 9 Gomila A, Carratalà J, Eliakim-Raz N, et al. Risk factors and prognosis of complicated urinary tract infections caused by Pseudomonas Aeruginosa in hospitalized patients: a retrospective multicenter cohort study. *Infect Drug Resist* 2018;11:2571–81.
- 10 Zorginstituut Nederland. Screeningsrapport Systematische analyse Infectieziekten. 2019. Available: https://www.zorginstituutnederland. nl/binaries/zinl/documenten/rapport/2019/05/14/zinnige-zorg--rapport-screeningsfase-infectieziekten/Zinnige+Zorg+-+Rapport+ screeningsfase+Systematische+analyse+Infectieziekten.pdf

- Bonkat G, Cai T, Veeratterapillay R, et al. Management of Urosepsis in 2018. Eur Urol Focus 2019;5:5–9.
- 12 World Health Organization. WHO methods and data sources for global burden of disease estimates 2000–2011. global health estimates technical paper. 2013. Available: http://www.who.int/healthinfo/statistics/GlobalDALYmethods_2000_2011.pdf
- 13 Haagsma JA, van Beeck EF, Polinder S, et al. The effect of Comorbidity on health-related quality of life for injury patients in the first year following injury: comparison of three Comorbidity adjustment approaches. *Popul Health Metrics* 2011;9.
- 14 Cassini A, Plachouras D, Eckmanns T, et al. Burden of six Healthcare-associated infections on European population health: estimating incidence-based disability-adjusted life years through a population prevalence-based Modelling study. *PLoS Med* 2016;13:e1002150.
- 15 van Lier A, de Gier B, McDonald SA, et al. Disease burden of Varicella versus other vaccine-preventable diseases before introduction of vaccination into the National Immunisation programme in the Netherlands. *Euro Surveill* 2019;24:1800363.
- 16 Murray CJL, Ezzati M, Flaxman AD, et al. GBD 2010: design, definitions, and Metrics. Lancet 2012;380:2063–6.
- 17 Colzani E, Cassini A, Lewandowski D, et al. A software tool for estimation of burden of infectious diseases in Europe using incidence-based disability adjusted life years. *PLoS One* 2017;12:e0170662.
- 18 Altorf-van der Kuil W, Schoffelen AF, de Greeff SC, et al. National laboratory-based surveillance system for antimicrobial resistance: a successful tool to support the control of antimicrobial resistance in the Netherlands. *Eurosurveillance* 2017;22.
- 19 Rijksinstituut Voor Volksgezondheid en milieu. Handleiding ISIS-AR, Bilthoven. 2017. Available: https://www.rivm.nl/sites/default/files/ 2018-11/HandleidingISIS-AR 2017 %28februari 2017%29.pdf
- 20 Rijksinstituut voor Volksgezondheid en Milieu. ISIS AR Populatie en Representativiteit. 2021. Available: https://www.rivm.nl/isis-ar/ populatie-en-representativiteit
- 21 Cai T, Palagin I, Brunelli R, et al. Office-based approach to urinary tract infections in 50 000 patients: results from the REWIND study. Int J Antimicrob Agents 2020;56:S0924-8579(20)30123-0.
- 22 ISTAT. Resident population by age, sex and marital status on 1ST January 2016 Italy. 2020. Available: http://demo.istat.it/pop2016/ index_e.html
- 23 Serretiello E, Folliero V, Santella B, et al. Trend of bacterial Uropathogens and their susceptibility pattern: study of single academic high-volume center in Italy (2015–2019). Int J Microbiol 2021;2021:5541706.
- 24 Cardone S, Petruzziello C, Migneco A, et al. Age-related trends in adults with urinary tract infections presenting to the emergency Department: A 5-year experience. *Rev Recent Clin Trials* 2019;14:147–56.
- 25 Luzzaro F, Ortisi G, Larosa M, et al. Prevalence and epidemiology of microbial pathogens causing bloodstream infections: results of the OASIS multicenter study. *Diagn Microbiol Infect Dis* 2011;69:363–9.
- 26 Tacconelli E, Pezzani MD. Public health burden of antimicrobial resistance in Europe. *Lancet Infect Dis* 2019;19:4–6.
- 27 Bordino V, Vicentini C, D'Ambrosio A, et al. Burden of Healthcareassociated infections in Italy: incidence, attributable mortality and disability-adjusted life years (Dalys) from a nationwide study, 2016. J Hosp Infect 2021;113:164–71.
- 28 Rottier WC, Deelen JWT, Caruana G, et al. Attributable mortality of antibiotic resistance in gram-negative infections in the Netherlands: a parallel matched cohort study. *Clin Microbiol Infect* 2020:S1198-743X(20)30420-1.
- 29 van Hout D, Verschuuren TD, Bruijning-Verhagen PCJ, et al. Extended-spectrum beta-Lactamase (ESBL)-Producing and non-ESBL-producing Escherichia coli isolates causing bacteremia in the Netherlands (2014–2016) differ in Clonal distribution, antimicrobial resistance gene and virulence gene content. *PLoS One* 2020;15:e0227604.
- 30 Collaborators AR. Articles global burden of bacterial antimicrobial resistance in 2019: a systematic analysis antimicrobial resistance collaborators. *Lancet* 2022:399.
- 31 Godijk NG, Bootsma MCJ, van Werkhoven HC, et al. Modelling addition and replacement mechanisms of Plasmid-based betalactam resistant E. coli infections. *Infectious Diseases (except HIV/ AIDS*) 2021.
- 32 Temkin E, Carmeli Y. Zero or more: methodological challenges of counting and estimating deaths related to antibiotic-resistant infections. *Clin Infect Dis* 2019;69:2029–34.
- 33 Pezzani MD, Tornimbene B, Pessoa-Silva C, et al. Methodological quality of studies evaluating the burden of drug-resistant infections in humans due to the WHO global antimicrobial

resistance surveillance system target bacteria. *Clin Microbiol Infect* 2021;27:687–96.

- 34 Lombardia F per i servizi degli O dei farmacisti della. Dispensazione Senza Ricetta: Quando SI Può E come SI FA. n.d. Available: https:// www.ordinifarmacistilombardia.it/farmacista/per_la_farmacia/ dispensazione_senza_ricetta.html?fbclid=IwAR2Hzk07wRFnygmyG5 Z1m4d5OkBnUbvFXqYMIfurvRbROzSms4XzU450PWM
- 35 Italian Medicines Agency. Antibiotics. n.d. Available: https://www. aifa.gov.it/en/farmaci-antibiotici?fbclid=lwAR2jIo2UTMVnOcHP80 us5MOjk9OpLwg21rYWWWi2Yvx7LdkKdusaVzqdKqs
- 36 MacKinnon MC, McEwen SA, Pearl DL, et al. Mortality in Escherichia coli bloodstream infections: a multinational population-based cohort study. BMC Infect Dis 2021;21:606.
- 37 Teslya A, Rozhnova G, Pham TM, et al. The importance of sustained compliance with physical distancing during COVID-19 vaccination Rollout. Commun Med (Lond) 2022;2:146. 10.1038/s43856-022-00207-3 Available: https://github.com/NoorGo/ExcessBurden
- 38 ECDC. Surveillance of antimicrobial resistance in Europe Annual report of the European Antimicrobial Resistance Surveillance Network (EARS-Net) 2017. Stockholm, 2018. Available: https://www. ecdc.europa.eu/sites/portal/files/documents/AMR-surveillance-EARS-Net-2017.pdf
- 39 Saint S. Clinical and economic consequences of Nosocomial catheter-related Bacteriuria. Am J Infect Control 2000;28:68–75.
- 40 Palacios-Baena ZR, Gutiérrez-Gutiérrez B, De Cueto M, *et al.* Development and validation of the INCREMENT-ESBL predictive score for mortality in patients with bloodstream infections due to

extended-spectrum-B-Lactamase-producing Enterobacteriaceae. J Antimicrob Chemother 2017;72:906–13.

- 41 Butler CC, Hillier S, Roberts Z, *et al.* Antibiotic-resistant infections in primary care are symptomatic for longer and increase workload: outcomes for patients with E. coli Utis. *Br J Gen Pract J R Coll Gen Pract* 2006;56:686–92.
- 42 Vallejo-Torres L, Pujol M, Shaw E, et al. Cost of hospitalised patients due to complicated urinary tract infections: a retrospective observational study in countries with high prevalence of multidrugresistant gram-negative bacteria: the COMBACTE-MAGNET, RESCUING study. BMJ Open 2018;8:e020251. 10.1136/ bmjopen-2017-020251 Available: http://bmjopen.bmj.com/content/8/ 4/e020251.abstract
- 43 Covino M, Manno A, Merra G, et al. Reduced utility of early Procalcitonin and blood culture determination in patients with febrile urinary tract infections in the emergency Department. Intern Emerg Med 2020;15:119–25.
- 44 de Kraker MEA, Wolkewitz M, Davey PG, et al. Burden of antimicrobial resistance in European hospitals: excess mortality and length of hospital stay associated with bloodstream infections due to Escherichia coli resistant to third-generation cephalosporins. J Antimicrob Chemother 2011;66:398–407.
- 45 Tumbarello M, Spanu T, Di Bidino R, et al. Costs of bloodstream infections caused by Escherichia coli and influence of extendedspectrum-beta-Lactamase production and inadequate initial antibiotic therapy. Antimicrob Agents Chemother 2010;54:4085–91.

Appendix 1. Systematic Review Methods

The Netherlands

PubMed and Embase were searched using the search terms shown in Appendix 3, resulting in 242 and 136 articles respectively. The removing of duplicates in Endnote and Rayyan resulted in a final set of 296 articles for title/abstract screening. In this stage, articles were included that reported Dutch studies on UTI or bacteraemia, that potentially contained data for both susceptible and resistant UTIs, but had not necessarily reported these data or did not mention the specific pathogen. In the case where, for example, testing for resistance had been mentioned but specific data were not separately reported for AMR and AMS *E. coli*, the authors were emailed. After full-text screening of 43 articles, a total of 18 were retained, and the authors were requested more data. If there was no response after a month, a follow-up email was sent.

We excluded case studies and studies that were carried out in a specific vulnerable population (elderly persons, children), or in highly-specific clinical patient populations. Inclusion criteria applied to the final set of articles were : a Dutch study, published in 2017 or later, UTI caused by resistant and/or susceptible *E. coli*, and estimates for of one or more of parameters needed for the OTs. Following this systematic literature search, further relevant articles were possibly identified during correspondence with authors.

Given the almost null yield of the first search, a second literature search was undertaken to locate relevant studies specifically informing the model parameters involving bacteraemia (i.e., P(Bact|UTI), DD(Bact), P(Death|Bact)) (Appendix 3). This produced 24 hits, due to the limited number of hits, we performed full-text screening for all. Inclusion criteria were only that the study reported suitable data on cases of bacteraemia in which *E. coli* had been isolated.

Following this search, further relevant articles were possibly identified in correspondence with authors of retained articles. We then applied the following algorithm to the set of identified articles: (i) if no eligible Dutch population studies were found reporting parameter values involving bacteraemia due to susceptible/resistant *E. coli* UTI, then (ii) Dutch studies reporting parameter values involving bacteraemia with susceptible/resistant *E. coli from any infection site* were used. (iii) If still no eligible studies found, then *EU* studies reporting parameter values due to bacteraemia with AMR/AMS *E. coli* from any infection site were considered eligible.

A third systematic literature search was conducted to attempt to find relevant studies specifically to inform P(Bact|UTI), with restriction to studies of resistant *E. coli* UTIs (Appendix 1). This produced 13 hits; 10 articles were eliminated based on abstract screening and the remaining three after full-text screening. A PRISMA diagram for all three searches together is shown in Figure S1.

Italy

PubMed and Embase were searched using the search term in Appendix 4, and yielded 231 and 176 results respectively. After removing duplicates in EndNote and Rayan, 290 articles remained. After title/abstract screening 56 articles were screened full text and 32 articles potentially contained parameter estimates relevant for the Italian population.

Given the almost null yield of the first search, we performed new separate searches for the incidence, progression from UTI to bacteraemia, DD(UTI) and LOS due to bacteraemia. For LOS(UTI) a third search was conducted (Appendix 4). Eventually, three articles from the search and one article recommended to the authors which fell outside the initial search criteria

of articles published from 2017 or later were used to estimate the parameters. A PRISMA

diagram for all searches on Italian parameters together is shown in Figure S2.

Appendix 2 – Systematic review to identity Dutch parameter estimates

Search 1

4th of February 2019

PubMed: (("2017/01/01"[Date - Publication] : "3000"[Date - Publication])) AND (((((urinary[Title/Abstract] AND tract[Title/Abstract]) AND (infection[Title/Abstract] OR infections[Title/Abstract])) OR urinary tract infection[MeSH] OR UTI[Title/Abstract]) AND (Netherlands OR Netherlands[MeSH] OR Dutch) AND (english[Language] OR dutch[Language])) AND ("2017/01/01"[PDat] : "3000/12/31"[PDat]))

242 results

4th of February 2019

Embase: ('urinary':ab,ti AND 'tract':ab,ti AND ('infection':ab,ti OR 'infections':ab,ti) OR 'uti':ab,ti) AND ('netherlands' OR 'dutch') AND [article]/lim AND ([dutch]/lim OR [english]/lim) AND [humans]/lim AND [embase]/lim AND [2017-2019]/py

136 results

Search 2

10th of February, 2020

PubMed: ((bacteraemia[Title/Abstract]) OR (bacteremia[Title/Abstract])) AND ((Netherlands[Text Word]) AND Dutch[Text Word]).

24 results

13th of February, 2020

PubMed:(((((urinary

tract[Title/Abstract])

AND infection[Title/Abstract]))

OR

UTI[Title/Abstract])) AND ((Netherlands[Text Word]) OR Dutch [Text Word])) AND ((lenght of stay[Text Word]) OR LOS[Text Word])

3 results

Search 3 –

13th of February, 2020

PubMed: (((bacteraemia[Text Word]) OR (bacteremia[Text Word])) AND resist*[Text Word] AND (E coli[MeSH] OR E coli[Text Word]) AND ((urinary tract infection[MeSH]) OR (UTI[Text Word])) AND (("probability of"[Text Word]) OR (progress*[Text Word]) OR ("risk of "[Text Word]))).

13 results



Figure S1

PRISMA flowchart of the first literature search on Dutch parameter estimates

Appendix 3 – Systematic review to identify Italian parameter estimates

Search 1

4th of February, 2019

Pubmed: ((("2017/01/01"[Date - Publication] : "3000"[Date - Publication])) AND (((((urinary[Title/Abstract] AND tract[Title/Abstract]) AND (infection[Title/Abstract] OR infections[Title/Abstract])) OR urinary tract infection[MeSH] OR UTI[Title/Abstract]) AND (Italy OR Italy[MeSH] OR Italian) AND (english[Language] OR dutch[Language])) AND ("2017/01/01"[PDat] : "3000/12/31"[PDat])))

231 results

('urinary':ab,ti AND 'tract':ab,ti AND ('infection':ab,ti OR 'infections':ab,ti) OR 'uti':ab,ti) AND ('italy' OR 'italian') AND [article]/lim AND ([dutch]/lim OR [english]/lim) AND [humans]/lim AND [embase]/lim AND [2017-2019]/py

176 results

Search 2

3th of June, 2020

Incidence - PubMed: ("2019/01/01"[Date - Publication] : "3000"[Date - Publication]) AND Italy AND ((urinary[Title/Abstract] AND tract [Title/Abstract] AND infection [Title/Abstract]) OR UTI [Title/Abstract]) AND incidence.

35 results

3th of June, 2020

LOS UTI - PubMed ("2019/01/01"[Date - Publication] : "3000"[Date - Publication]) AND Italy AND ((urinary[Title/Abstract] AND tract [Title/Abstract] AND infection [Title/Abstract]) OR UTI [Title/Abstract]) AND (LOS [Title/Abstract] OR (length [Title/Abstract] AND stay [Title/Abstract])).

5 results

18th of June, 2020

UTI to bacteraemia - PubMed: (("2019/01/01"[Date - Publication] : "2020/06/18"[Date -

Publication])) AND ((Italy[Text Word]) AND ((UTI[Title/Abstract]) OR

(((urinary[Title/Abstract]) AND (tract[Title/Abstract])) AND (infection[Title/Abstract])))).

21 results

31st of August, 2020

 LOS Bacteraemia – PubMed: (("2005/01/01"[Date - Publication] : "3000"[Date - Publication]))

 AND ((Italy[Text Word])
 AND (((((bacteraemia[Title/Abstract]))
 OR

 (bacteraemias[Title/Abstract]))
 OR (bacteremia[Title/Abstract]))
 OR

 (bacteremias[Title/Abstract]))
 OR (((length[Title/Abstract]))
 OR

 (bacteremias[Title/Abstract]))
 AND ((((days[Title/Abstract]))
 OR (((length[Title/Abstract]))

 AND (of[Title/Abstract]))
 AND (stay[Title/Abstract])))
 OR (LOS[Title/Abstract]))))

24 results

Search 3

16th of June, 2020

LOS UTI- Pubmed (("2015/01/01"[Date - Publication] : "3000"[Date - Publication]) AND Italy AND ((urinary[Title/Abstract] AND tract [Title/Abstract] AND infection [Title/Abstract]) OR UTI [Title/Abstract]) AND (LOS [Title/Abstract] OR disability duration [Title/Abstract] OR (length [Title/Abstract] AND stay [Title/Abstract]))) NOT (("2019/01/01"[Date - Publication] : "2020/06/02"[Date - Publication]) AND Italy AND ((urinary[Title/Abstract] AND tract [Title/Abstract] AND infection [Title/Abstract]) OR UTI [Title/Abstract]) AND (LOS [Title/Abstract] OR (length [Title/Abstract] AND stay [Title/Abstract]))).

7 results

2nd of September, 2020

(((general practitioner) OR (general practice)) AND (((urinary tract infection) OR (UTI))

AND (Italy))) AND ((out-patient) OR (outpatient))

10 results, 1 included



Figure S2

PRISMA flowchart of the literature search on Italian parameter estimates



Figure S3

smoothed geographical distribution of the percentage of inhabitants for whom at least 1 urinary isolate was found in the ISIS-AR database in 2018, by 4-digit postal code area and with regional cooperative network borders

Appendix 4.

Systematic review results

The Netherlands

The first systematic literature review yielded only two articles, both providing an estimate of DD(UTI). In the first study the Netherlands was one of four countries on which analysis was based, and bacteria species and AMS vs. AMR infections were not distinguished. Correspondence with the authors yielded a more appropriate citation (1), which was carried out in England and Wales in 2002-2004 and reported DD(UTI) for *E. coli* UTIs separately for AMS and AMR infection. (In Figure S1 this article is indicated as an additional record identified through other sources). We justified this choice as the analysis in (2) did not find any between-country difference in DD(UTI).

The second review resulted in one suitable study for P(Death|Bact), which was a Dutch study that reported 30-day mortality in bacteraemia patients with either resistant *E. coli* or susceptible *E. coli* in 2014–2016 (3). The study population had a median age of 69 years (*IQR* 57 to 77); it is plausible that a lower mortality rate would be observed for younger age-groups. We could only locate a single study reporting age-group specific values for 30-day mortality due to bacteraemia (4). This study was conducted in Iceland among patients with bacteraemia caused by *S. aureus*. We took the simple approach of setting the parameter values for P(Death|Bact) for the age-groups 55 years and older to the value from study (3), and then scaling the parameter values for the younger age-group, 55-74 years, and the <35 years and 35-54 years age-groups from the Icelandic study (4). This meant P(Death|Bact) was zero for <35

years (since mortality risk was 0% for <35 years (4)), and a scaling factor of 0.54 (from 3.8%/7.1%) was applied to 35-54 years.

For the parameter DD(Bact), the literature reviews did not yield any eligible studies. We decided to adopt values from (5), which is a large well-conducted multi-country study that was carried out in 2007/8, and that reported patient characteristic-adjusted LOS values for both AMS and 3rd generation cephalosporin-resistant *E. coli* bloodstream infections (BSIs). All selected parameter values are provided in Table 1.

As the third systematic literature review, which was specifically aimed at P(Bact|UTI), did not yield any studies. We relied on a previous pooled analysis [24] which we identified through citation search. This study did not distinguish between AMR and AMS infections, and the contributing studies were all carried out in the USA in the 1980s.

Italy

We found one article providing estimates on P(Death|Bact). The study of Palacios-Baena et al. (6) found a 30-day mortality of 26.2% of ESBL blood stream infections (BSI), 34 of the 130 Italian BSI patients died. We calculated the mortality for susceptible BSI using the ratio of susceptible vs. resistant mortality reported in another, less recent, Italian study by Tumbarello et al. (7) and estimated a 30-day mortality of 5.47% for susceptible BSI.

Furthermore, for DD(UTI) and DD(BACT) we only found an Italian study amongst elderly (Mdn = 77, IQR = 65-83) with UTIs or urosepsis which reported a mean LOS of 10 [7-17] days (8) and a median LOS of 9.5 days for Italian patients with complicated UTIs in Italy (9). Of the UTIs 58% was caused by *E. coli*. Unfortunately, no studies were identified which specified LOS for ESBL *E. coli* and *E. coli* UTIs. Because the lack of better studies on DD(UTI) and

DD(BACT) amongst adults, we used the estimate of Covino et al. (8) in elderly and Vallejo-Torres et al. (9) on complicated UTIs.

Moreover, we searched the citations of Cassini et al. (10) for relevant Italian studies and found that Tumbarello et al. (7) reported LOS for resistant BSI of 20 ± 17 days and 13 ± 9 days for non-AMR BSI.

For P(Bact | UTI) we were unable to locate a parameter and, therefore, we used the same value as the Dutch parameter. For the health outcomes following bacteremia, other than death, we used the same values as Cassini et al. (10).

Regarding the incidence of resistant *E. coli*, we did not locate any direct estimates; therefore, we estimated incidence (see Methods).

References Appendix 4.

- Butler CC, Hillier S, Roberts Z, Dunstan F, Howard A, Palmer S. Antibiotic-resistant infections in primary care are symptomatic for longer and increase workload: outcomes for patients with E. coli UTIs. Br J Gen Pract J R Coll Gen Pract. 2006 Sep;56(530):686–92.
- Altorf-van der Kuil W, Schoffelen AF, de Greeff SC, Thijsen SF, Alblas HJ, Notermans DW, et al. National laboratory-based surveillance system for antimicrobial resistance: a successful tool to support the control of antimicrobial resistance in the Netherlands. Euro Surveill Bull Eur sur les Mal Transm = Eur Commun Dis Bull. 2017 Nov;22(46).
- 3. van Hout D, Verschuuren TD, Bruijning-Verhagen PCJ, Bosch T, Schürch AC,
 Willems RJL, et al. Extended-spectrum beta-lactamase (ESBL)-producing and nonESBL-producing Escherichia coli isolates causing bacteremia in the Netherlands (2014 2016) differ in clonal distribution, antimicrobial resistance gene and virulence gene
 content. PLoS One [Internet]. 2020 Jan 14;15(1):e0227604. Available from:
 https://doi.org/10.1371/journal.pone.0227604
- Asgeirsson H, Gudlaugsson O, Kristinsson KG, Heiddal S, Kristjansson M. Staphylococcus aureus bacteraemia in Iceland, 1995-2008: changing incidence and mortality. Clin Microbiol Infect Off Publ Eur Soc Clin Microbiol Infect Dis. 2011 Apr;17(4):513–8.
- 5. de Kraker MEA, Wolkewitz M, Davey PG, Koller W, Berger J, Nagler J, et al. Burden of antimicrobial resistance in European hospitals: excess mortality and length of hospital stay associated with bloodstream infections due to Escherichia coli resistant

to third-generation cephalosporins. J Antimicrob Chemother. 2011 Feb;66(2):398–407.

- 6. Palacios-Baena ZR, Gutiérrez-Gutiérrez B, De Cueto M, Viale P, Venditti M, Hernández-Torres A, et al. Development and validation of the INCREMENT-ESBL predictive score for mortality in patients with bloodstream infections due to extended-spectrum-β-lactamase-producing Enterobacteriaceae. J Antimicrob Chemother. 2017 Mar;72(3):906–13.
- Tumbarello M, Spanu T, Di Bidino R, Marchetti M, Ruggeri M, Trecarichi EM, et al. Costs of bloodstream infections caused by Escherichia coli and influence of extendedspectrum-beta-lactamase production and inadequate initial antibiotic therapy. Antimicrob Agents Chemother [Internet]. 2010/07/26. 2010 Oct;54(10):4085–91. Available from: https://pubmed.ncbi.nlm.nih.gov/20660675
- Covino M, Manno A, Merra G, Simeoni B, Piccioni A, Carbone L, et al. Reduced utility of early procalcitonin and blood culture determination in patients with febrile urinary tract infections in the emergency department. Intern Emerg Med. 2020 Jan;15(1):119–25.
- 9. Vallejo-Torres L, Pujol M, Shaw E, Wiegand I, Vigo JM, Stoddart M, et al. Cost of hospitalised patients due to complicated urinary tract infections: a retrospective observational study in countries with high prevalence of multidrug-resistant Gramnegative bacteria: the COMBACTE-MAGNET, RESCUING study. BMJ Open [Internet]. 2018 Apr 1;8(4):e020251. Available from: http://bmjopen.bmj.com/content/8/4/e020251.abstract
- 10. Cassini A, Högberg LD, Plachouras D, Quattrocchi A, Hoxha A, Simonsen GS, et al.

Attributable deaths and disability-adjusted life-years caused by infections with antibiotic-resistant bacteria in the EU and the European Economic Area in 2015: a population-level modelling analysis. Lancet Infect Dis [Internet]. 2019 Jan 1;19(1):56– 66. Available from: https://doi.org/10.1016/S1473-3099(18)30605-4

Table S1. Number and incidence of resistant E. coli UTI per age- and sex category in the Netherlands in 2018

			Males					Females		
					Average					
				Recurrent*	resistant				Recurrent*	Average
		Number of	Resistant	resistant	E. coli		Number of	Resistant	resistant	resistant
Age	Male	resistant	E.coli UTIs	E. Coli	UTIs per	Female	resistant	E. coli UTIs	E. coli	E. coli UTIs
category	inhabitants	E. coli UTIs	incidence	UTIs	patient	inhabitants	E. coli UTIs	incidence	UTIs	per patient
0	87001	12	0.000137929	0	1.00	82565	10	0.000121117	0	1.00
1-4	358019	25	5.86561E-05	4	1.19	340514	117	0.000323041	7	1.06
5-9	475503	19	2.10304E-05	9	1.90	452563	148	0.000287253	18	1.14
10-14	494511	11	1.61776E-05	3	1.38	471948	62	0.000122895	4	1.07
15-19	536852	20	2.79407E-05	5	1.33	511180	60	0.000105638	6	1.11
20-24	542817	19	2.76336E-05	4	1.27	525964	140	0.000230054	19	1.16
25-29	560319	36	5.53256E-05	5	1.16	545838	155	0.000283967	0	1.00
30-34	530554	38	6.59688E-05	3	1.09	522235	136	0.000250845	5	1.04
35-39	512925	26	3.70425E-05	7	1.37	512431	114	0.000204906	9	1.09

40-44	516723	53	6.77346E-05	18	1.51	521589	125	0.000191722	25	1.25
45-49	634188	92	0.000108801	23	1.33	634635	198	0.000272598	25	1.14
50-54	644223	143	0.000176957	29	1.25	635623	270	0.00035713	43	1.19
55-59	606130	211	0.000268919	48	1.29	605380	441	0.000597972	79	1.22
60-64	537540	287	0.000401831	71	1.33	542198	445	0.000671341	81	1.22
65-69	495875	460	0.000703806	111	1.32	503662	473	0.000770358	85	1.22
70-74	424486	633	0.001036548	193	1.44	447439	613	0.001115236	114	1.23
75-79	273902	621	0.001595461	184	1.42	314838	689	0.001715168	149	1.28
80-84	172825	487	0.002065673	130	1.36	235430	683	0.002229962	158	1.30
≥85	122648	495	0.003000457	127	1.35	248011	1056	0.00033063	236	1.29
Total	8527041	3688	0.000432506	974	1.36	8654043	5935	0.000685807	1063	1.22

*Defined as a UTI occurring more than 14 days after another UTI

Table S2. Number and incidence of E. coli UTI per age- and sex category in the Netherlands in 2018

			Males					Females		
					Average				Recurrent	
Age	Male		E.coli	Recurrent	E. coli				*	Average
categor	inhabitant	Number of	UTIs	* E. Coli	UTIs per	Female	Number of	E. coli UTIs	E. coli	E. coli UTIs
у	S	E. coli UTIs	incidence	UTIs	patient	inhabitants	E. coli UTIs	incidence	UTIs	per patient
0	87001	453	0.0052	0	1.00	82565	413	0.0044	52	1.14
1-4	358019	598	0.0015	74	1.14	340514	4079	0.0105	518	1.15
5-9	475503	351	0.0006	45	1.15	452563	6336	0.0115	1111	1.21
10-14	494511	260	0.0005	34	1.15	471948	2766	0.0049	473	1.21
15-19	536852	315	0.0005	41	1.15	511180	2651	0.0047	260	1.11
20-24	542817	318	0.0005	30	1.10	525964	3499	0.0061	316	1.10
25-29	560319	492	0.0008	60	1.14	545838	3745	0.0069	0	1.00
30-34	530554	500	0.0009	25	1.05	522235	3714	0.0069	102	1.03
35-39	512925	731	0.0012	106	1.17	512431	3638	0.0063	432	1.13
40-44	516723	968	0.0016	147	1.18	521589	3608	0.0060	497	1.16

Total	8527041	49528	0.0058	9006	1.22	8654043	149913	0.0173	25177	1.20
≥85	122648	6251	0.0411	1213	1.24	248011	22890	0.0747	4355	1.23
80-84	172825	6148	0.0280	1312	1.27	235430	16251	0.0548	3340	1.26
75-79	273902	7017	0.0203	1465	1.26	314838	15964	0.0407	3162	1.25
70-74	424486	7930	0.0149	1598	1.25	447439	16474	0.0297	3201	1.24
65-69	495875	6022	0.0099	1113	1.23	503662	12863	0.0210	2296	1.22
60-64	537540	4166	0.0065	672	1.19	542198	10635	0.0161	1880	1.21
55-59	606130	3154	0.0044	498	1.19	605380	8686	0.0120	1418	1.20
50-54	644223	2224	0.0029	331	1.17	635623	6648	0.0089	1022	1.18
45-49	634188	1630	0.0022	242	1.17	634635	5053	0.0068	742	1.17

*Defined as a UTI occurring more than 14 days after another UTI

Table S3. Sensitivity analysis of the number and incidence of resistant E. coli UTI per age- and sex category in the Netherlands in 2018

			Males					Females		
					Average					
				Recurrent*	resistant				Recurrent*	Average
		Number of	Resistant	resistant	E. coli		Number of	Resistant	resistant	resistant
Age	Male	resistant	E.coli UTIs	E. Coli	UTIs per	Female	resistant	E. coli UTIs	E. coli	E. coli UTIs
category	inhabitants	E. coli UTIs	incidence	UTIs	patient	inhabitants	E. coli UTIs	incidence	UTIs	per patient
0	87001	12	0.000137929	0	1.00	82565	10	0.000121117	0	1.00
1-4	358019	21	5.86561E-05	0	1.00	340514	111	0.000323041	1	1.01
5-9	475503	10	2.10304E-05	0	1.00	452563	137	0.000287253	7	1.05
10-14	494511	8	1.61776E-05	0	1.00	471948	58	0.000122895	0	1.00
15-19	536852	16	2.79407E-05	1	1.07	511180	54	0.000105638	0	1.00
20-24	542817	16	2.76336E-05	1	1.07	525964	121	0.000230054	0	1.00
25-29	560319	32	5.53256E-05	1	1.03	545838	155	0.000283967	0	1.00
30-34	530554	35	6.59688E-05	0	1.00	522235	132	0.000250845	1	1.01
35-39	512925	19	3.70425E-05	0	1.00	512431	107	0.000204906	2	1.02

Total	8527041	2908	0.000341	194	1.07	8654043	5111	0.000591	239	1.05
≥85	122648	400	0.003000457	32	1.09	248011	879	0.003306305	59	1.07
80-84	172825	379	0.002065673	22	1.06	235430	563	0.002229962	38	1.07
75-79	273902	477	0.001595461	40	1.09	314838	581	0.001715168	41	1.08
70-74	424486	488	0.001036548	48	1.11	447439	526	0.001115236	27	1.05
65-69	495875	368	0.000703806	19	1.05	503662	409	0.000770358	21	1.05
60-64	537540	227	0.000401831	11	1.05	542198	382	0.000671341	18	1.05
55-59	606130	169	0.000268919	6	1.04	605380	377	0.000597972	15	1.04
50-54	644223	118	0.000176957	4	1.04	635623	231	0.00035713	4	1.02
45-49	634188	75	0.000108801	6	1.09	634635	178	0.000272598	5	1.03
40-44	516723	38	6.77346E-05	3	1.09	521589	100	0.000191722	0	1.00

*Defined as a UTI occurring more than 3 months after another UTI