

Clinical predictors of escalating care in hepatic and renal cyst infection in autosomal dominant polycystic kidney and liver disease

M.A. Lantinga¹, R.G.L. de Sévaux², T.J.G. Gevers¹, W.J.G. Oyen³, J.W. de Fijter⁴,
D. Soonawala⁴, R. Zietse⁵, M. Salih⁵, N.F. Casteleijn⁶, E.M. Spithoven⁷, E. Meijer⁷,
R.T. Gansevoort⁷, J.P.H. Drenth^{1*}; on behalf of the DIPA consortium

Departments of ¹Gastroenterology and Hepatology, ²Nephrology, ³Nuclear Medicine, Radboud University Medical Center, Nijmegen, the Netherlands, ⁴Department of Nephrology, Leiden University Medical Center, Leiden, the Netherlands, ⁵Department of Nephrology, Erasmus Medical Center Rotterdam, Rotterdam, the Netherlands, ⁶Department of Urology, University Medical Center Groningen, University of Groningen, Groningen, the Netherlands, ⁷Department of Nephrology, University Medical Center Groningen, University of Groningen, Groningen, the Netherlands, *corresponding author: e-mail: joostphdrenth@cs.com.

ABSTRACT

Background: Cyst infection may occur in autosomal dominant polycystic kidney disease (ADPKD) and autosomal dominant polycystic liver disease (ADPLD). Antimicrobial agents often fail to control infection, leading to invasive action. We aimed to identify factors predicting escalation of care.

Methods: ADPKD and ADPLD patients were identified from local/national databases (2001-2013). Records were screened for patients meeting criteria for cyst infection (positive cyst aspirate and/or clinical findings). Factors that predict escalated care were identified with multivariate modified Poisson regression.

Results: We screened 1773 patients. A total of 77 patients with cyst infection (4.3%) were included for analysis (hepatic 36%; male 49%; age 54 ± 13 years; ADPKD 95%; dialysis 9%, diabetes 18%, renal transplant 56%, eGFR [IQR 24-78] ml/min/1.73 m² (excluding patients with a history of renal transplant or receiving dialysis)). A pathogen was identified in 71% of cases. *Escherichia coli* was the most common pathogen and accounted for 69% of cases. Initial treatment was limited to antibiotics in 87% of patients (n = 67), 40% included a fluoroquinolone. Ultimately, 48% of patients underwent some form of invasive action (escalation of care). Increasing white blood cell count (WBC) (RR 1.04 95%-CI 1.01-1.07, p = 0.008) was associated with escalating care, whereas an increase in time between transplant and infection (RR 0.92 95% CI 0.86-0.97, p = 0.005) and *E. coli* isolation (RR 0.55 95% CI 0.34-0.89, p = 0.02) were protective.

Conclusion: High serum WBC, isolation of atypical pathogens and early infection after transplantation are factors that increase the risk of escalation of care in hepatic and renal cyst infection patients.

KEYWORDS

ADPKD, ADPLD, cyst infection

INTRODUCTION

Cyst infection may occur as a complication in autosomal dominant polycystic kidney disease (ADPKD) and autosomal dominant polycystic liver disease (ADPLD) and results in considerable morbidity and mortality.^{1,2} Cyst infections are thought to arise from haematogenous spread of translocated gut bacteria (hepatic cyst infection) or ascending urinary tract infection (renal cyst infection).³ *Escherichia coli* is the most frequently isolated microorganism in both hepatic and renal cyst infection.⁴ Diagnosing cyst infection may be difficult as a definite diagnosis can only be made upon presence of cyst aspirate with inflammatory cells and bacteria.⁵ In most cases, a cyst aspirate is not routinely taken. Therefore, physicians are likely to make clinical decisions using a mix of clinical, biochemical and imaging findings,⁶ which led to the development of several diagnostic algorithms for patients in whom cyst infection is suspected.^{7,8}

The first line of treatment of cyst infection consists of antibiotics. There is no evidence base for the choice of antibiotic therapy. Fluoroquinolones are traditionally selected due to their favourable pharmacokinetic properties as ciprofloxacin readily diffuses in hepatic and renal cysts.^{9,10} However, antibiotics often fail to control the infection. This leads to frequent switching between antibiotic classes and ultimately to invasive action in an attempt to control the infection.¹¹

We hypothesise that patient-related and clinical factors (age, gender, serum inflammation parameters, comorbidities, history of solid organ transplantation) modify the risk for additional invasive measures that are required to control the infection.¹² The aim of the present study was to investigate factors that predict escalation of care in a nationwide cohort of 77 patients who developed cyst infection.

MATERIALS AND METHODS

Ethical consideration

This study did not require formal review by the Institutional Review Board (IRB) after study protocol evaluation by the local IRB (CMO Radboud University Medical Center Nijmegen, registration number 2013/299).

Study design

We established a multicentre retrospective cohort of patients who were diagnosed with cyst infection. Patients were enrolled at four tertiary referral centres in the Netherlands which participated in the DIPAK-1 Study. This cohort study is reported in accordance with the Strengthening The Reporting of OBServational studies in Epidemiology (STROBE) statement (*supplementary table 1**).¹³

Setting and participants

We identified ADPKD and ADPLD patients through inspection of a number of local and national databases. The local databases are established because of financial reimbursement systems that register the diagnosis of patients using general Diagnosis Treatment Combinations (DTCs). The national database consisted of RENINE (Registration Renal Replacement Therapy the Netherlands), a Dutch foundation that registers all renal transplant recipients in the Netherlands. The DIPAK-1 Study group is an investigator-driven, multicentre, open label, randomised controlled trial, planned to enrol 300 individuals who are aged 18-60 years, with an estimated glomerular filtration rate (eGFR) of 30-60 ml/min/1.73m².¹⁴

We included ADPKD and ADPLD patients diagnosed with either hepatic or renal cyst infection. We diagnosed

cyst infection in case of a positive cyst aspirate (white blood cells and bacteria) or when cyst infection was considered by the individual physician to be stated in the medical record in the combined presence of: 1) serum C-reactive protein (CRP) > 50 mg/l, 2) body temperature $\geq 38.5^{\circ}\text{C}$ and 3) abdominal pain.^{6,15} Medical records were screened for patients who met the inclusion criteria for cyst infection. We excluded patients if: a) cyst infection was a complication of aspiration sclerotherapy, b) the site of cyst infection was not documented, c) data on initial treatment were incomplete or d) ADPLD or ADPKD was not reported in the medical chart.

Variables

We extracted patient demographics and clinical variables at the time of cyst infection from medical records into an electronic database. The variables identified included gender, age at cyst infection, type of polycystic disease, site of cyst infection, peak serum CRP, peak serum white blood cell count (WBC), peak estimated glomerular filtration rate (eGFR), microbiological culture results (urine, blood and cyst aspirate), initial treatment, initial antibiotic treatment, treatment duration and details on invasive treatment. We also collected data on comorbidities including history of solid organ transplant, interval between solid organ transplant and cyst infection, dialysis, diabetes mellitus and immunosuppressive therapy. Definitions of variables are discussed in more detail in *supplementary table 2*.

Outcomes

We defined any procedure that aimed to breach the wall of the infected cyst, either through percutaneous access or surgery as escalation of care.

Multiple imputation

We used multiple imputation to account for missing data. We limited imputation of missing variables to patients in whom the outcome was available.¹⁶ Multiple imputation relies on the assumption that data are missing at random, which implicates that the probability that a specific variable is missing depends on the other variables that are available for that patient (i.e. other observed patient characteristics).¹⁷ Literature suggests to impute at least 40 times to allow accurate estimation of missing variables.¹⁶

Analysis

We performed all statistical analyses using SPSS statistical software package version 22 (SPSS Inc., Chicago, IL). Chi-square test was used for dichotomous data, Student t-test for parametric data and Mann-Whitney U test for non-parametric data. All tests were two-tailed and a p value of < 0.05 was considered statistically significant.

As the expected incidences of the outcomes investigated in this study were high, we calculated adjusted risk ratios

to avoid any possible misinterpretation of odds ratios (OR) as risk ratios (RR).¹⁸ We selected Poisson regression as an alternative for binary logistic regression to obtain adjusted RR instead of OR.¹⁹ To correct for the use of binary outcomes, we used a robust error variance procedure to correct for the incorrect assumption of Poisson distributed outcomes.²⁰

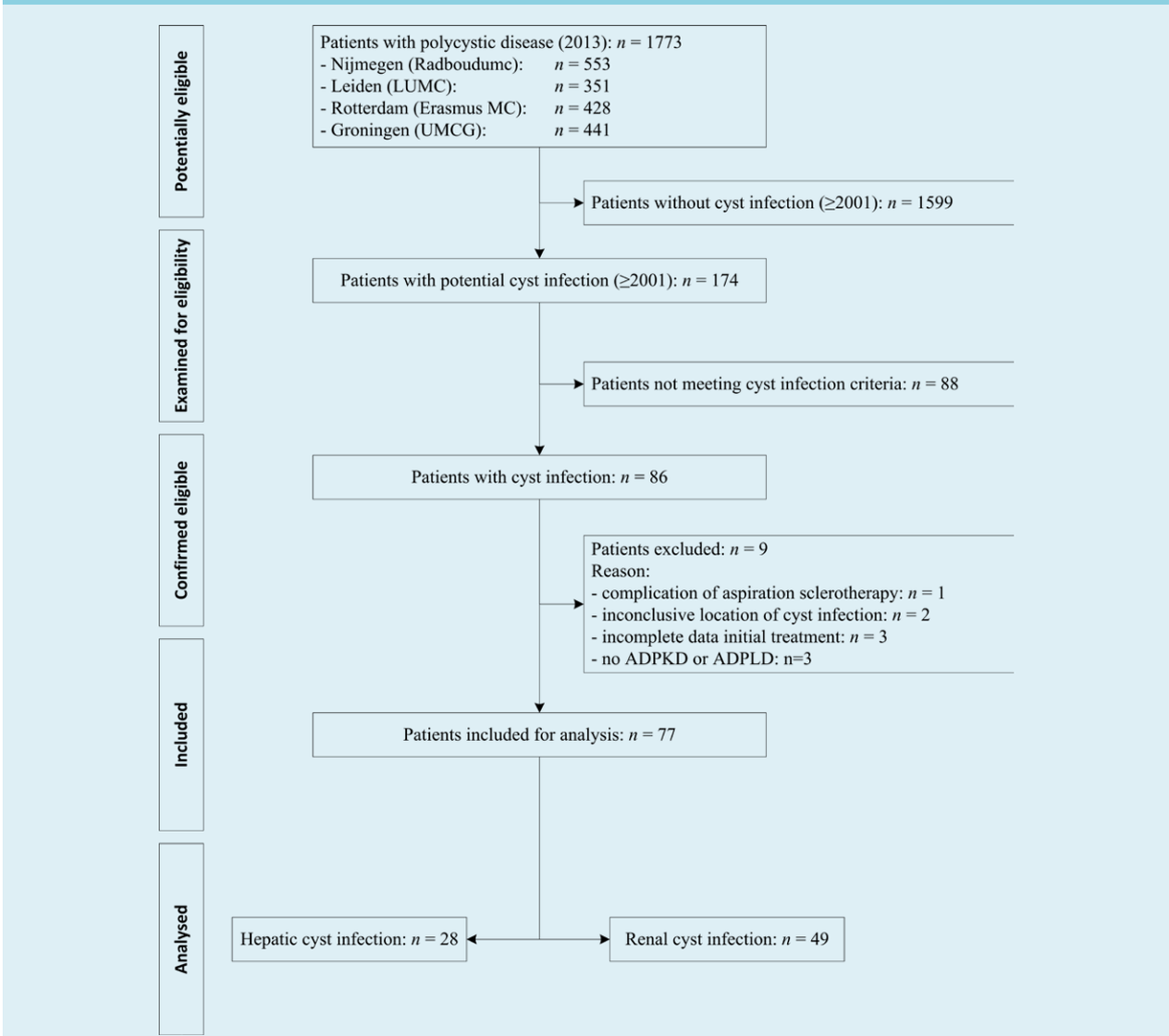
Variables with a p value of < 0.2 in univariate analysis were included in the multivariate modified Poisson regression model with manual backward elimination of non-significant variables. A p value of < 0.05 was considered to be statistically significant. We performed univariate and multivariate analyses on both the original and imputed datasets.

RESULTS

Cyst infection patients

We identified 1773 polycystic disease patients in four Dutch tertiary referral centres (figure 1). We examined 174 patients (9.8%) for eligibility and 86 (4.9%) had developed hepatic or renal cyst infection in or after 2001. Patients were excluded if the infection developed following aspiration sclerotherapy (n = 1), infection location was unknown (n = 2), data on initial treatment were missing (n = 3) or polycystic disease was absent (n = 3). Ultimately, we included 77 cyst infection patients (4.3%) for analysis.

Figure 1. Number of patients at each stage of the study



Radboudumc = Radboud University Medical Center; UMCG = University Medical Center Groningen; LUMC = Leiden University Medical Center; Erasmus MC = Erasmus Medical Center Rotterdam; ADPKD = autosomal dominant polycystic kidney disease; ADPLD = autosomal dominant polycystic liver disease; n = number

Patient demographics

Table 1 shows characteristics of the included patients with cyst infection (n = 77). Almost all patients suffered from ADPKD (95%). Of these patients, 49% were male and the average age at infection was 54 years. Hepatic cyst infection was diagnosed in 28 patients (36%), whereas

49 patients (64%) were diagnosed with a renal cyst infection. Diabetes mellitus was frequent (18%) and 56% of patients with cyst infection had a history of solid organ transplantation (all kidney transplant recipients). The median interval between transplantation and cyst infection was four years.

Table 1. Characteristics extracted from medical records of cyst infection patients

Characteristic	Total (n = 77)	Missing (n)
Male, n (%)	38 (49)	
Age at cyst infection, years (\pm SD)	54 \pm 13	
Type of polycystic disease, n (%)		
- ADPKD	73 (95)	
- ADPLD	4 (5)	
Site of cyst infection, n (%)		
- Liver	28 (36)	
- Kidney	49 (64)	
Diabetes mellitus, n (%)	14 (18)	
Dialysis, n (%)	7 (9)	
Solid organ transplant, n (%) ^a	43 (56)	
- Interval solid organ transplant and cyst infection, years [IQR]	4 [1-10]	
Immunosuppressive drug use, n (%)	44 (57)	
Peak serum CRP, mg/l [IQR]	215 [123-283]	1
Peak serum WBC, $\times 10^9/l$ [IQR]	11.6 [8.3-15.7]	1
Peak eGFR, ml/min/1.73m ² [IQR] ^b		6
- MDRD	50 [24-78]	
- CKD-EPI	57 [26-87]	
Positive microbiological culture, n (%) ^c	55 (71)	
- Hepatic cyst infection patients	23 (82)	
- Renal cyst infection patients	32 (65)	
Atypical pathogen cultured (other than <i>E. coli</i>), n (%)	24 (31)	3
Initial treatment		
- Limited to antibiotics, n (%) ^d	67 (87)	
- Containing fluoroquinolone, n (%)	27 (40)	1
- Duration, days [IQR]	8 [4-39]	6
Follow-up, years [IQR]	2 [1-5]	

Parametric variables are expressed as mean \pm SD. Non-parametric variables are expressed as median [IQR]. Percentages may not add up to 100 due to rounding.

n = number; SD = standard deviation; IQR = interquartile range; ADPKD = autosomal dominant polycystic kidney disease; ADPLD = autosomal dominant polycystic liver disease; CRP = C-reactive protein; WBC = white blood cell count; MDRD = modification of diet in renal disease; CKD-epi = Chronic Kidney Disease Epidemiology Collaboration; *E.coli* = *Escherichia coli*; N/A = not applicable.

^a All involved kidney transplantations.

^b Excluding patients with a history of renal transplant or actively receiving dialysis.

^c Supplementary table 4 provides an overview of isolated pathogens.

^d Table 2 provides an overview of antibiotic regimens.

Follow-up

Median time of follow-up was two years [IQR 1-5 years]. Follow-up duration was similar between hepatic and renal cyst infection patients (1 year [IQR 0-5 years] vs. 3 years [IQR 1-6 years], $p = 0.06$) (supplementary table 3).

Characteristics at time of diagnosis

Peak serum CRP and white blood cell count were clearly raised, reaching a median level of 215 mg/l (upper limit of normal (ULN): < 10 mg/l) and $11.6 \times 10^9/l$ (ULN: < $11 \times 10^9/l$), respectively. In most patients renal function was estimated to be mildly impaired (MDRD: 50 ml/min/1.73 m², CKD-epi: 57 ml/min/1.73 m²). *E. coli* was isolated in most urine (57%) and blood (66%) cultures (supplementary table 4). A substantial proportion of cultures returned alternative pathogens.

Treatment characteristics

Initial treatment was limited to the use of antibiotics in 87% ($n = 67$) of cyst infection patients; the remainder of the patients received antibiotics in combination with invasive treatment. Some 40% ($n = 67$) received a fluoroquinolone antibiotic, either as mono or combination antibiotic treatment (table 1). The median duration of initial treatment was eight days. Table 2 provides an overview of all antibiotics given as initial treatment.

Differences between hepatic and renal cyst infection patients

Characteristics of hepatic and renal cyst infection patients were comparable, except for age (58 years \pm 10 vs. 51 years \pm 14, $p = 0.02$) and peak serum WBC ($9.4 \times 10^9/l$ [IQR 7.8-12.3] vs. $12.8 \times 10^9/l$ [IQR 9.2-18.5], $p = 0.007$) (supplementary table 3).

Escalation of care during follow-up

In 48% of patients ($n = 32$) escalation of care occurred (table 3, supplementary table 5). The peak serum WBC appeared to be significantly higher in these patients ($12.5 \times 10^9/l$ [IQR 9.1-18.5] vs. $10.9 \times 10^9/l$ [IQR 8.3-13.8], $p = 0.01$). Furthermore, atypical pathogens were more frequent (47% vs. 21%, $p = 0.01$). In contrast, the interval between transplantation and cyst infection was significantly shorter in patients receiving invasive treatment compared with those who did not (3 years [IQR 0-8] vs. 9 years [IQR 1-14], $p = 0.03$). Follow-up duration was comparable between patients regardless of escalation of care (2 years [IQR 1-6] vs. 2 years [IQR 1-5], $p = 0.61$).

Predictors for escalation of care

Selected variables (p value < 0.2 in univariate analysis) were entered into the multivariate model (table 4). Multiple imputation had no effect on the selection of predictors (supplementary table 6). Exclusive isolation of *E. coli*

Table 2. Antibiotic regimens in patients with initial treatment limited to antibiotics

Antibiotic regimen	Initial cyst infection treatment limited to antibiotics (n = 67)
Monotherapy, n (%)	56 (84)
Fluoroquinolones, n (%)	26 (46)
- Ciprofloxacin, %	100
Cephalosporins, n (%)	16 (29)
2nd generation	
- Cefuroxime, %	56
3rd generation	
- Ceftriaxone, %	19
- Ceftazidime, %	19
- Cefotaxime, %	6
Penicillin, n (%)	9 (16)
- Amoxicillin/clavulanic acid, %	78
- Amoxicillin, %	11
- Piperacillin/tazobactam, %	11
Carbapenems, n (%)	3 (5)
- Meropenem, %	100
Glycopeptides, n (%)	1 (2)
- Vancomycin, %	100
Sulphonamides, n (%)	1 (2)
- Trimethoprim/sulfamethoxazole, %	100
Combination therapy, n (%)	10 (15)
- Cefuroxime + gentamicin, %	20
- Cefuroxime + tobramycin, %	20
- Cefuroxime + metronidazole, %	10
- Cefazolin + metronidazole, %	10
- Amoxicillin + teicoplanin, %	10
- Ceftazidime + vancomycin, %	10
- Ciprofloxacin + gentamicin, %	10
- Flucloxacillin + ceftazidime, %	10
Missing, n (%)	1 (1)

Percentages may not add up to 100 due to rounding.

decreased the risk for escalation of care (RR 0.55, 95% CI 0.34-0.89, $p = 0.02$) (figure 2). In contrast, if peak serum WBC increased or the interval between transplantation and cyst infection decreased, patients were more likely to be exposed to escalation of care (RR 1.04, 95% CI

Table 3. Univariate modified Poisson regression to identify predictors for escalation of care during follow-up.

Variable			Original dataset	Imputed datasets
	Conservative (n = 45)	Escalated care (n = 32) ^a	P value	P value
Gender, n (%)				
- Male	21 (47)	17 (53)	0.58	0.58
- Female (referent)	24 (53)	15 (47)		
Age at cyst infection, years (± SD)	51 ± 13	57 ± 13	0.06	0.06
Type of polycystic disease, n (%)				
- ADPLD	2 (4)	2 (6)	0.71	0.71
- ADPKD (referent)	43 (96)	30 (94)		
Site of cyst infection, n (%)				
- Liver	18 (40)	10 (31)	0.44	0.44
- Kidney (referent)	27 (60)	22 (69)		
Diabetes mellitus, n (%)				
- No	37 (82)	26 (81)	0.91	0.91
- Yes (referent)	8 (18)	6 (19)		
Dialysis, n (%)				
- No	41 (91)	29 (91)	0.94	0.94
- Yes (referent)	4 (9)	3 (9)		
Solid organ transplant, n (%)^b				
- No	22 (49)	12 (38)	0.33	0.33
- Yes (referent)	23 (51)	20 (62)		
Interval solid organ transplant and cyst infection, years [IQR]	9 [1-14]	3 [0-8]	0.02	0.03
Immunosuppressive drug use, n (%)				
- No	22 (49)	11 (34)	0.22	0.22
- Yes (referent)	23 (51)	21 (66)		
Peak serum CRP, mg/l [IQR]	194 [126-269]	228 [122-306]	0.22	0.32
Peak serum WBC, $\times 10^9/l$ [IQR]	10.9 [8.3-13.8]	12.5 [9.1-18.5]	0.002	0.01
Peak eGFR, ml/min/1.73m² [IQR]^c				
- MDRD	45 [27-71]	66 [17-88]	0.56	0.55
- CKD-EPI	52 [28-81]	70 [17-95]	0.65	0.67
Atypical pathogen cultured, n (%)				
- No	33 (79)	17 (53)	0.02	0.01
- Yes (referent)	9 (21)	15 (47)		
Initial antibiotic treatment containing fluoroquinolone, n (%)				
- No	26 (58)	16 (53)	0.70	0.68
- Yes (referent)	19 (42)	14 (47)		
Duration initial antibiotic treatment, days [IQR]	9 [5-42]	7 [4-39]	0.55	0.70
Follow-up, years	2 [1-6]	2 [1-5]	0.61	0.61

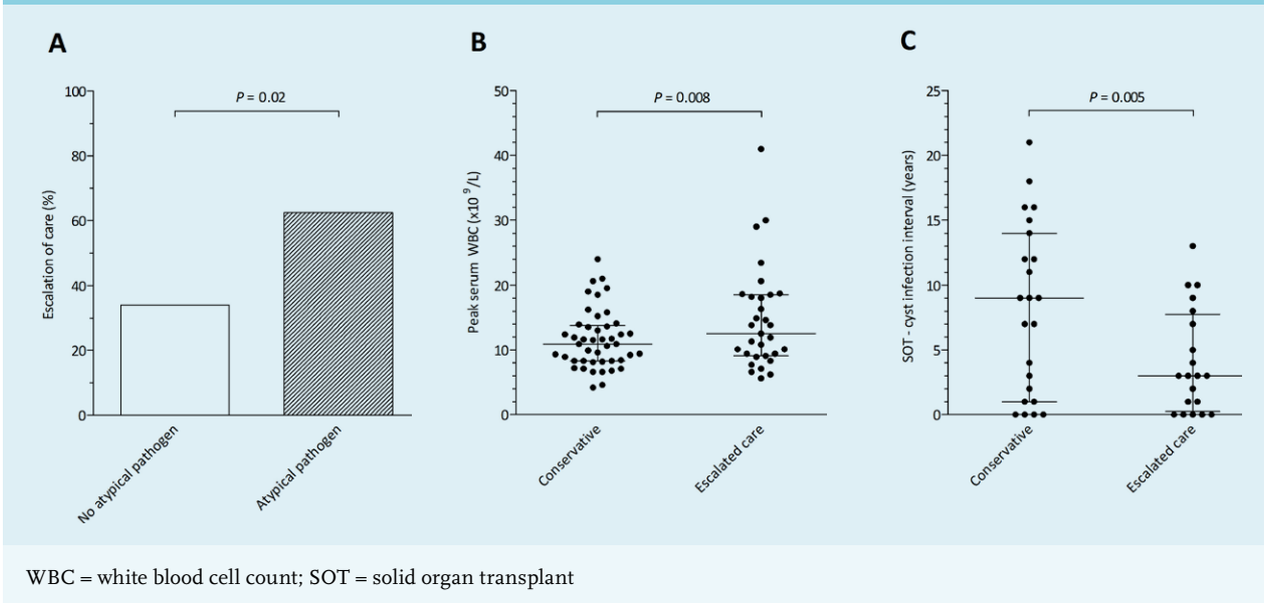
Percentages may not add up to 100 due to rounding.
CI = confidence interval; ADPKD = autosomal dominant polycystic kidney disease = ADPLD = autosomal dominant polycystic liver disease; CRP = C-reactive protein; WBC = white blood cell count; MDRD = modification of diet in renal disease; CKD-epi = Chronic Kidney Disease Epidemiology Collaboration.
^a Supplementary table 5 shows details on escalated care.
^b All involved kidney transplants.
^c Excluding patients with history of renal transplant or actively receiving dialysis.

Table 4. Multivariate modified Poisson regression to identify predictors for escalation of care during follow-up (imputed datasets).

Variable	Poisson model before backward elimination of non-significant variables		Poisson model after backward elimination of non-significant variables	
	Multivariate adjusted risk ratio (95% CI)	P value	Multivariate adjusted risk ratio (95% CI)	P value
Age at cyst infection (years)	1.02 (0.99-1.04)	0.14		
Interval solid organ transplant and cyst infection (years)	0.92 (0.87-0.97)	0.003	0.92 (0.86-0.97)	0.005
Peak serum WBC ($\times 10^9/l$)	1.04 (1.01-1.07)	0.006	1.04 (1.01-1.07)	0.008
Atypical pathogen cultured				
- No	0.63 (0.39-0.99)	0.05	0.55 (0.34-0.89)	0.02
- Yes (referent)				

CI = confidence interval; WBC = white blood cell count.

Figure 2. Rate of escalation of care according to (A) cultured pathogen, (B) peak serum white blood cell count and (C) interval between solid organ transplant and cyst infection



1.01-1.07, $p = 0.008$ and RR 0.92, 95% CI 0.86-0.97, $p = 0.005$, respectively). Age at cyst infection was discarded in the final model. In contrast to the imputed datasets, the presence or absence of atypical pathogens did not predict escalation of care in the final model using the original dataset (supplementary table 7).

DISCUSSION

The key finding of our multicentre retrospective cohort study is that increasing serum WBC, isolation of pathogens other than *E. coli* and a shorter transplant-infection

interval increases the risk for escalating care to the point of puncture or operation of the cyst.

We identified several factors in our cohort that increase the risk for treatment escalation. Previous studies on cyst infection involved single-centre cohort studies and focused on the clinical, microbiological and imaging aspects of cyst infection. None specifically investigated factors that influence the need for invasive treatment.^{3-4,15,21-23} In this study we found that patients with a longer interval between solid organ transplantation and development of cyst infection were less likely to be exposed to escalation of care. This observation might be explained by a reduced need for immunosuppressive drugs in patients who are

clinically stable.^{24,25} This could explain the observation that these patients are more capable of clearing the infection. Alternatively, the early period after solid organ transplantation could warrant a more aggressive (i.e. invasive) approach to cyst infection to benefit the outcome of transplanted allografts.^{26,27}

Increasing serum WBC and isolation of atypical pathogens both independently increase the risk for treatment escalation. Serum WBC reflects infection severity. In a study investigating acute pyelonephritis, a severe increase in serum WBC predicted clinical failure.²⁸ *E. coli* is isolated in most cases of cyst infection.⁶ In the presence of an alternative pathogen, treatment limited to antibiotics could be less effective as successful eradication of alternative bacteria might be more challenging. Two systematic reviews on the management of hepatic and renal cyst infection, respectively, show that in case an alternative micro-organism is isolated (e.g. *Enterococcus faecium*), escalation of care is more likely.^{11,29}

In patients with high serum WBC, isolation of atypical pathogens or a history of solid organ transplantation, the risk of failing conventional antibiotic treatment increases. Given these results, it is reasonable to consider alternative treatment options such as percutaneous cyst drainage, (partial) resection or even kidney and/or liver transplantation.

The main strength of this study comes with its size as we identified ADPKD and ADPLD patients who developed hepatic or renal cyst infection in four Dutch tertiary referral centres. This approach, combined with the use of broad inclusion criteria, enhances the generalisability of our findings. A potential limitation is the retrospective nature of our study and use of data not primarily intended for study purposes. This resulted in the potential exclusion of a large proportion of patients who did have cyst infection but did not meet the inclusion criteria due to missing data. We used multiple imputation to correct for missing data and, as such, reduced bias and loss of power. Moreover, the modified Poisson regression model is very reliable even in relatively small sample sizes.²⁰ We did not include total liver volume and total kidney volume as predictors. A recent study showed that infectious complications in ADPKD do not correlate with the total liver volume;³⁰ it remains unclear whether total kidney volume predicts cyst infection outcome. Finally, we combined patients with hepatic and renal cyst infection. Univariate analysis showed an imbalance for age and peak serum WBC (*supplementary table 5*). We hypothesised that its effect on the outcome would be limited as both covariates have a theoretical effect on the outcomes in both hepatic and renal cyst infection patients. We corrected for this by including these covariates in the multivariate model. Moreover,

adding the site of cyst infection to the final multivariate model had no significant effect (*supplementary table 8*).

To conclude, we show that high serum WBC, presence of pathogens other than *E. coli* and early infection after transplantation predict the need for change to invasive procedures.

*The supplementary material is available on request from the editorial office (marina@alphatekst.nl).

ACKNOWLEDGEMENTS

The authors wish to thank Wietske Kievit and Marcia Tummers from the Department of Health Evidence, Radboud University Nijmegen, the Netherlands for their methodological and statistical help. Furthermore, the authors wish to thank Aline Hemke, coordinator of RENINE.

DIPAK Consortium

The DIPAK Consortium is an inter-university collaboration in the Netherlands that is established to study autosomal dominant polycystic kidney disease and to develop rational treatment strategies for this disease. The DIPAK Consortium is sponsored by the Dutch Kidney Foundation (grant CP10.12). Principal investigators are (in alphabetical order): J.P.H. Drenth (Department of Gastroenterology and Hepatology, Radboud University Medical Center Nijmegen), J.W. de Fijter (Department of Nephrology, Leiden University Medical Center), R.T. Gansevoort (Department of Nephrology, University Medical Center Groningen), D.J.M. Peters (Department of Human Genetics, Leiden University Medical Center), J. Wetzels (Department of Nephrology, Radboud University Medical Center Nijmegen), R. Zietse (Department of Internal Medicine, Erasmus Medical Center Rotterdam).

DISCLOSURES

Prof. Dr. Drenth has received grant support and fees for serving on advisory boards and consultancy from IPSEN and Novartis. Prof. Dr. Gansevoort received grant support and fees for serving on advisory boards and steering committees from IPSEN, Otsuka Pharmaceuticals and Sanofi-Genzyme.

No other potential conflict of interest relevant to this article was reported.

No grant support was received for this study/article.

REFERENCES

1. Gevers TJ, Drenth JP. Diagnosis and management of polycystic liver disease. *Nat Rev Gastroenterol Hepatol*. 2013;10:101-8.
2. Torres VE, Harris PC, Pirson Y. Autosomal dominant polycystic kidney disease. *Lancet*. 2007;369:1287-301.
3. Suwabe T, Araoka H, Ubara Y, et al. Cyst infection in autosomal dominant polycystic kidney disease: causative microorganisms and susceptibility to lipid-soluble antibiotics. *Eur J Clin Microbiol Infect Dis*. 2015;34:1369-79.
4. Jouret F, Lhommel R, Beguin C, et al. Positron-emission computed tomography in cyst infection diagnosis in patients with autosomal dominant polycystic kidney disease. *Clin J Am Soc Nephrol*. 2011;6:1644-50.
5. Jouret F, Lhommel R, Devuyt O, et al. Diagnosis of cyst infection in patients with autosomal dominant polycystic kidney disease: attributes and limitations of the current modalities. *Nephrol Dial Transplant*. 2012;27:3746-51.
6. Lantinga MA, Drenth JP, Gevers TJ. Diagnostic criteria in renal and hepatic cyst infection. *Nephrol Dial Transplant*. 2015;30:744-51.
7. Neuville M, Hustinx R, Jacques J, Krzesinski JM, Jouret F. Diagnostic Algorithm in the Management of Acute Febrile Abdomen in Patients with Autosomal Dominant Polycystic Kidney Disease. *PLoS One*. 2016;11(8):e0161277.
8. Lantinga MA, Darding AJM, de Sevaux RGL, et al. International Multi-Specialty Delphi Survey: Identification of Diagnostic Criteria for Hepatic and Renal Cyst Infection. *Nephron*. 2016;134:205-14.
9. Hiyama L, Tang A, Miller LG. Levofloxacin penetration into a renal cyst in a patient with autosomal dominant polycystic kidney disease. *Am J Kidney Dis*. 2006;47(1):e9-13.
10. Telenti A, Torres VE, Gross JB, Jr., et al. Hepatic cyst infection in autosomal dominant polycystic kidney disease. *Mayo Clin Proc*. 1990;65:933-42.
11. Lantinga MA, Geudens A, Gevers TJ, Drenth JP. Systematic review: the management of hepatic cyst infection. *Aliment Pharmacol Ther*. 2015;41:253-61.
12. Chapman AB, Devuyt O, Eckardt KU, et al. Autosomal-dominant polycystic kidney disease (ADPKD): executive summary from a Kidney Disease: Improving Global Outcomes (KDIGO) Controversies Conference. *Kidney Int*. 2015;88:17-27.
13. von Elm E, Altman DG, Egger M, et al. The Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement: guidelines for reporting observational studies. *Lancet*. 2007;370:1453-7.
14. Meijer E, Drenth JP, d'Agnolo H, et al. Rationale and design of the DIPAK 1 study: a randomized controlled clinical trial assessing the efficacy of lanreotide to Halt disease progression in autosomal dominant polycystic kidney disease. *Am J Kidney Dis*. 2014;63:446-55.
15. Sallee M, Rafat C, Zahar JR, et al. Cyst infections in patients with autosomal dominant polycystic kidney disease. *Clin J Am Soc Nephrol*. 2009;4:1183-9.
16. Graham JW, Olchowski AE, Gilreath TD. How many imputations are really needed? Some practical clarifications of multiple imputation theory. *Prev Sci*. 2007;8:206-13.
17. Donders AR, van der Heijden GJ, Stijnen T, Moons KG. Review: a gentle introduction to imputation of missing values. *J Clin Epidemiol*. 2006;59:1087-91.
18. Knol MJ. [Down with odds ratios: risk ratios in cohort studies and randomised clinical trials]. *Ned Tijdschr Geneesk*. 2012;156:A4775.
19. Knol MJ, Le Cessie S, Algra A, Vandenbroucke JP, Groenwold RH. Overestimation of risk ratios by odds ratios in trials and cohort studies: alternatives to logistic regression. *CMAJ*. 2012;184:895-9.
20. Zou G. A modified poisson regression approach to prospective studies with binary data. *Am J Epidemiol*. 2004;159:702-6.
21. Bobot M, Ghez C, Gondouin B, et al. Diagnostic performance of [(18)F]fluorodeoxyglucose positron emission tomography-computed tomography in cyst infection in patients with autosomal dominant polycystic kidney disease. *Clin Microbiol Infect*. 2016;22:71-7.
22. Suwabe T, Ubara Y, Higa Y, et al. Infected hepatic and renal cysts: differential impact on outcome in autosomal dominant polycystic kidney disease. *Nephron Clin Pract*. 2009;112:c157-63.
23. Suwabe T, Ubara Y, Sumida K, et al. Clinical features of cyst infection and hemorrhage in ADPKD: new diagnostic criteria. *Clin Exp Nephrol*. 2012;16:892-902.
24. Zand MS. Immunosuppression and immune monitoring after renal transplantation. *Semin Dial*. 2005;18:511-9.
25. Kirk AD, Mannon RB, Swanson SJ, Hale DA. Strategies for minimizing immunosuppression in kidney transplantation. *Transpl Int*. 2005;18:2-14.
26. Cainelli F, Vento S. Infections and solid organ transplant rejection: a cause-and-effect relationship? *Lancet Infect Dis*. 2002;2:539-49.
27. Chong AS, Alegre ML. The impact of infection and tissue damage in solid-organ transplantation. *Nat Rev Immunol*. 2012;12:459-71.
28. Wie SH, Ki M, Kim J, et al. Clinical characteristics predicting early clinical failure after 72 h of antibiotic treatment in women with community-onset acute pyelonephritis: a prospective multicentre study. *Clin Microbiol Infect*. 2014;20:O721-9.
29. Lantinga MA, Casteleijn NF, Geudens A, et al. Management of renal cyst infection in patients with autosomal dominant polycystic kidney disease: a systematic review. *Nephrol Dial Transplant*. 2017;32:144-50.
30. Kim H, Park HC, Ryu H, et al. Clinical Correlates of Mass Effect in Autosomal Dominant Polycystic Kidney Disease. *PLoS One*. 2015;10:e0144526.