

# Left ventricular assist device-related infections: a multicentric study

S. Simeon, E. Flecher, M. Revest, M. Niculescu, J. -C. Roussel, M. Michel, P.

Leprince, P. Tattevin

## ► To cite this version:

S. Simeon, E. Flecher, M. Revest, M. Niculescu, J. -C. Roussel, et al.. Left ventricular assist device-related infections: a multicentric study. Clinical Microbiology and Infection, 2017, 23 (10), pp.748-751. 10.1016/j.cmi.2017.03.008 . hal-01619296

# HAL Id: hal-01619296 https://univ-rennes.hal.science/hal-01619296

Submitted on 7 Jun2018

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#### 1 Original article

2 Left ventricular assist device-related infections: a multicentric study

## 3 Running title: Left ventricular assist device-related infections

- 4 S. Siméon<sup>1</sup>, E. Flécher<sup>2</sup>, M. Revest<sup>1,3</sup>, M. Niculescu<sup>4</sup>, J-C Roussel<sup>5</sup>, M.
- 5 Michel <sup>6</sup>, P. Leprince <sup>7</sup>, P. Tattevin  $^{1,3,*}$
- 6 1) Department of Infectious Diseases and Intensive Care Unit, Pontchaillou University
- 7 Hospital, Rennes, France
- 8 2) Department of Cardio-Thoracic and Vascular Surgery, Pontchaillou University Hospital,
- 9 Rennes, France
- 10 3) Inserm U835, Rennes-1 University, France
- 4) Anesthesiology Department, Institut de Cardiologie, Groupe Hospitalier Pitié-Salpêtrière,
- 12 Université Pierre et Marie Curie, Assistance Publique des Hôpitaux de Paris, France
- 13 5) Department of Cardio-Thoracic and Vascular Surgery, Thorax Institute, Laennec
- 14 University Hospital, Nantes, France
- 15 6) Department of Cardiovascular Diseases, Laennec University Hospital, Nantes, France
- 16 7) Surgery Department, Institut de Cardiologie, Groupe Hospitalier Pitié-Salpêtrière,
- 17 Université Pierre et Marie Curie, Assistance Publique des Hôpitaux de Paris, France
- 18 \* Corresponding Author: P. Tattevin, Infectious Diseases and Intensive Care Unit,
- 19 Pontchaillou University Hospital, 2, rue Henri Le Guilloux, 35033 Rennes Cedex, France.
- 20 Tel +33 299289564. Fax +33 299282452 E-mail address: pierre.tattevin@chu-rennes.fr

#### 21 ABSTRACT

Objectives. Implantable left ventricular assist device (LVAD) is a major therapeutic progress for end-stage heart failure in selected patients. As their use is expanding, infectious complications are emerging, with limited data available to guide their management. We aimed to better characterize LVAD-related infections.

Methods. We enrolled all consecutive patients diagnosed with LVAD-related infections in 3
referral centers in France, using standardized definition of infections in patients with LVAD.
Data were collected from medical charts using a standardized questionnaire.

Results. Between 2007 and 2012, 159 patients received LVAD for end-stage heart failure. 29 Among them, 36 (22.6%) –5 women, 31 men– presented at least one infectious complication, 30 after a median time of 2.9 months from LVAD implantation [interquartile range, 1.8-7.5], 31 with a median follow-up of 12 months [8-17]. Main comorbidities were alcoholism (33%), 32 33 diabetes (11%) and immunosuppression (11%). Mean age at implantation was 51 ( $\pm$ 11) years. LVAD were implanted as bridge-to-transplant (n=22), bridge-to-recovery (n=8), destination 34 35 therapy (n=4), or unspecified (n=2). LVAD-related infections were restricted to the driveline exit site (n=17), had loco-regional extension (n=13), or reached the internal pump (n=3). The 36 main bacteria isolated were *Staphylococcus aureus* (n=20), coagulase-negative staphylococci 37 (n=7), Enterobacteriaceae (n=14), Pseudomonas aeruginosa (n=10) and Corynebacterium 38 sp. (n=7), with polymicrobial infections in 19 cases. LVAD could be retained in all patients, 39 with the use of prolonged antibacterial treatment in 34 (94%), and debridement in 17 (47%). 40 One patient died due to LVAD-associated infection. 41

42 Conclusions. LVAD-related infections are common after LVAD implantation, and may be
43 controlled by prolonged antibiotic treatment.

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#### 45 Introduction

Heart failure is a major cause of morbidity and mortality worldwide [1,2]. The prevalence and incidence of heart failure are increasing [3,4], and the constant shortage of donor organs increases the need for alternatives to heart transplant in patients with end-stage heart failure refractory to medical treatment [5]. Currently, around 2% of the adult population in developed countries suffers from heart failure [6].

In this context, the advent of implantable left ventricular assist device (LVAD) 51 represents a major medical progress for end-stage heart failure in selected patients [7,8], and 52 53 is currently used as a bridge-to-heart transplantation, a bridge-to-recovery, or as destination therapy (i.e., as the last resort in patients with neither perspectives of recovery, nor heart 54 transplant). Implantable LVAD intended for long-term use rely on a percutaneous driveline, 55 to carry electric signals and energy from the controller and batteries to the implanted pump. 56 As with any other implantable foreign device, LVAD is subject to LVAD-related infections, 57 a consequence of medical progress that is gradually emerging, proportionally to the number 58 of patients implanted with LVAD [9,10]. Indeed, the presence of a driveline piercing the skin 59 places the patient at a continuous risk for infections, that can affect the exit site, the 60 subcutaneous tunnel, the abdominal pocket (if present), the implanted pump, and disseminate 61 through bloodstream infections. The transition from pulsatile to continuous-flow LVAD 62 significantly improved clinical outcome [11], and decreased the risk of infectious 63 complications, but LVAD-related infections are still common [12,13]. Due to the scarcity of 64 data currently available in the medical literature, the management of these emerging 65 infections is poorly standardized, and mostly derives from the state-of-the-art for the 66 management of other cardiovascular devices-related infections (e.g. pacemaker, intra-cardiac 67 defibrillator, prosthetic valves, or vascular prosthesis), although their characteristics are 68

significantly different. We aimed to better characterize LVAD-related infections, their
treatment, and their outcome, through a multi-center study in three referral centers in France.

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#### 73 Methods

La Pitié-Salpêtrière is a 1663-bed university hospital located in Paris (France); Laennec is a 74 489-bed university hospital located in Nantes (Pays-de-Loire, France); and Pontchaillou is a 75 992-bed university hospital located in Rennes (Bretagne, France). They all serve as referral 76 centers for end-stage heart failure in their area. Although no national guidelines are currently 77 available for LVAD implantation in France, all three centers use antibacterial prophylaxis 78 79 with cefamandole for less than 24 hours from the time of LVAD implantation, under rigorous aseptic conditions, with no continuous antibacterial prophylaxis following implantation. 80 Throughout the study period, skin preparation procedures included preoperative shower with 81 chlorhexidine gluconate solution the night before surgery, and two separate skin preparation 82 before incision with either povidone-iodine or chlorhexidine with ethanol in the three 83 participating sites. 84

Patients with LVAD receive repeated counselling and education by specialized nurse 85 before and after LVAD implantation, to reduce the risk of infection and trauma at the exit 86 site. When LVAD-related complication is suspected, cases are reviewed by the endocarditis 87 team in each site, including at least one cardiac surgeon, one infectious diseases specialist, 88 and one microbiologist. In the absence of any consensus on the management of LVAD-89 related infections during the study period, patients were managed on a case-by-case basis, 90 taking into account clinical and microbiological data, including drug susceptibility profiles. 91 Follow-up was tailored to the characteristics of LVAD-related complications, and was mostly 92 performed by the cardiac surgery department, in association with infectious diseases 93

94 specialist for any suspicion of infectious complication. All patients enrolled in these95 databases provided informed consent for observational studies.

Cases of LVAD-related infections were identified through a retrospective review of 96 97 medical charts for all patients with LVAD implanted from January 2007 to December 2012 in the participating centers. LVAD-related infections were defined according to criteria 98 established by the Interagency Registry for Mechanically Assisted Circulatory Support 99 (INTERMACS) [14], the North American registry for mechanical circulatory support devices 100 (MCSD) that serves as a quality improvement system to assess the characteristics, treatments, 101 and outcomes of patients receiving MCSD. Briefly, percutaneous site infections were defined 102 as pain, erythema, or purulent drainage restricted to the LVAD entry site, with a positive 103 104 culture from the skin, and the decision to initiate systemic antimicrobial therapy. LVADrelated infections were defined as loco-regional when the erythema or induration extended >1 105 cm along the subcutaneous part of the driveline, and/or in case of fever or leukocytosis not 106 explained by other conditions, with the decision to initiate systemic antimicrobial therapy. 107 Lastly, LVAD-related infections where classified as pump infections when an 108 indistinguishable organism (genus, species, and antimicrobial susceptibility pattern) was 109 recovered from 2 or more peripheral blood cultures taken at least 12 hours apart with no other 110 focus of infection, leading to the initiation of an antimicrobial treatment. 111

A standardized questionnaire was used to collect demographical, clinical, and laboratory data from medical records, with a focus on the clinical features and microbiology of LVAD-related infections, any surgical and antibacterial treatment, and outcome. Statistical analyses were descriptive. Categorical variables were presented as number and percentages. Continuous variables were presented as either means with standard deviations (SD), or medians with first- and third-quartile [interquartile range, IQR], if the distribution of the data

118 were skewed. Statistical analyses were performed using STATA software, version 12119 (STATA Corporation).

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122 **Results** 

Between January 2007 and December 2012, 159 patients underwent LVAD implantation at 123 La Pitié-Salpêtrière, Paris (n=103), Laennec, Nantes (n=32), and Pontchaillou, Rennes 124 (n=24). Of these, 36 patients (22.6%) presented at least one LVAD-related infection during 125 the study period, with a median follow-up of 12 months [IQR, 8-17]. Patients characteristics 126 and features of LVAD-related infections are presented in tables 1 and 2. The median delay 127 from LVAD implantation to the diagnosis of LVAD-related infection was 2.9 months [1.8-128 7.5]. Microbiology documentation of LVAD-related infections was obtained using swabs 129 taken from the entry site (56%), surgical samples during debridement (20%) or during heart 130 transplant (7%), blood culture (10%), and transcutaneous puncture (3%). Infections were 131 polymicrobial in 54% of cases. Main pathogens were Staphyloccocus aureus (n=20, 132 including 2 meticillin-resistant), Enterobacteriaceae (n=12, including 7 extended spectrum 133 producers), *Pseudomonas* aeruginosa beta-lactamase (n=10), coagulase-negative 134 staphylococci (n=6, including 2 meticillin-resistant), Corynebacterium sp. (n=6), and others 135 (Acinetobacter baumanii, Haemophilus aprophilus, and Stenotrophomonas malthophilia, one 136 137 patient each).

LVAD-related infections required hospital admission in 28 cases (78%). All patients received systemic antibacterial treatment, based on pathogen(s) identification and drug susceptibility testing. Antibacterial treatment was temporarily administered by the intravenous route in 27 patients (75%), while 9 patients only received oral antibiotics. The median total duration of antibacterial treatment for LVAD-related infections was 90 days [40-

143 120] per patient. Surgical debridement and drainage were performed in 17 patients (47%). 144 Indications for surgery were either loco-regional infection associated with collection 145 documented by computed tomography scanning or ultrasound (n=12), or pump infection 146 (n=5). Procedures were always implemented with concomitant antibacterial treatment. Three 147 of the 17 patients treated with debridement required a second local surgery due to persistent 148 drainage on appropriate antibacterial treatment. No patient required wound vacuum-assisted-149 closure (VAC), pump relocation, or antibiotic beads at the device pocket.

The outcome was satisfactory in most cases: LVAD was finally extracted because of 150 heart transplant (n=17) - with a median time to transplantation of 11.3 months [8.9-15.0], or 151 because of recovery from heart failure (n=4). No patient underwent LVAD extraction 152 because of uncontrolled infection, and no LVAD-related infection was considered as a 153 definitive contra-indication for heart transplant. During the study period, 3 patients died, of 154 whom one was considered as attributable to LVAD-related infection. Of the 12 patients with 155 the LVAD still in place by the time of last follow-up, three had stopped all antibacterial 156 treatment - with a follow-up of 52, 98, and 138 days since last antibacterial treatment - and 157 nine were on prolonged oral antibacterial treatment. Most patients (n=34, 94%) presented no 158 sequel of their LVAD-related infection. 159

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162 **Discussion** 

163 The most salient features of this multi-center study of LVAD-related infections 164 performed in three referral centers are as follows: Firstly, we confirmed that infectious 165 complications are very common early after LVAD implantation, documented in almost one 166 quarter of patients, with a median follow-up of one year. This is in agreement with previous

167 series, and not unexpected given the continuous percutaneous portal of entry, and the comorbidities presented by these patients with end-stage heart failure. Secondly, we provided 168 a comprehensive documentation of the major pathogens involved in this emerging foreign 169 170 device-related infectious diseases, where the 'big five' are - as could be expected - S. aureus, Enterobacteriaceae, P. aeruginosa, coagulase-negative staphylococci, and Corynebacterium 171 sp. Thirdly, we found that, despite the major challenges arising when infections occur on a 172 life-saving device, the overall outcome was favorable in most instances, at the price of 173 prolonged systemic antibacterial treatment for > 90% of patients, and surgical debridement 174 for almost half of them. This satisfactory outcome may be related to the destiny of these 175 devices, mostly implanted as bridge-to-transplantation, or bridge-to-recovery: Indeed, in this 176 situation, the main challenge for the management of LVAD-related infections is to keep the 177 device in place until it is no longer necessary, which is not the case for other foreign device-178 related infections, such as prosthetic valve, vascular prosthesis, or prosthetic joints, usually 179 planned to remain in place lifelong. 180

To our knowledge, few studies have reported the incidence, the characteristics, and 181 the management of LVAD-related infections thus far. Previous reports found an incidence of 182 LVAD-related infections similar to our cohort, but with a median time from LVAD 183 implantation that was slightly longer, at 4.4 to 7.4 months [13,15], as compared to 2.9 months 184 in our study. These discrepancies may be related to the close monitoring routinely 185 implemented after LVAD implantation in our centers, to limited sample size, or to the fact 186 187 that 58% of our patients were classified as INTERMACS I by the time of implantation, which has been associated with earlier first pump-related infection [16]. Microbiological 188 characteristics of previous series are remarkably similar to ours, although one previous series 189 also isolated fungi as potentially involved (17). Of note, 22 isolates were Gram negative 190 bacilli in our series, including 19 non-susceptible to the antibioprophylactic regimen routinely 191

192 used during LVAD implantation in our sites (i.e. cefamandole): extended spectrum betalactamase producing Enterobacteriaceae (n=7), and non-fermeting Gram negative bacilli 193 (n=12). However, given the median delay from LVAD implantation to infection diagnosis 194 195 (2.9 months), this is unlikely that an antibiopropylactic regimen with broader coverage would have prevented these infections. We found that percutaneous driveline exit is the most 196 common site of LVAD-related infections, as in previous reports. Given that a number of 197 studies have demonstrated that trauma at the exit site favors the development of driveline 198 infections [10,18], prevention of driveline trauma may decrease the risk of LVAD-related 199 infections. Hence, patient education is an important component for the prevention of LVAD-200 related infections, and goes behind hygiene and infection control education. 201

Up to now, the management of LVAD-related infections is poorly standardized, due 202 to the scarcity of therapeutic studies in the field, and the absence of consensus guidelines. To 203 assist therapeutic decisions in this challenging context, Koval et al proposed 204 recommendations based on observational studies and experts opinion [18]. The management 205 of LVAD-related infections in our center follows the main principles depicted in these 206 recommendations, including anti-infectious treatment adapted to microbiology in all cases, 207 with surgical revision in selected cases. Although LVAD exchange is an option, this high-risk 208 surgery is often contra-indicated, and the use of continuous or iterative course of antibiotics 209 until transplantation, or recovery, is the rule for most centers. 210

Our study has limitations. First, although data were prospectively collected, the original database did not focus on infectious complications. Hence, collection of clinical data and microbiological investigations were not standardized. To address this limitation, the three participating sites have implemented a prospective database for all patients who receive LVAD since January 2012, with systematic follow-up at the referral center at one month and three months after implantation, and then at least every 6 months, as well as each time

217 requested by the patient and/or his general practitioner. Second, due to limited sample size, and the participation of three centers with similar practices in a restricted geographic area, 218 our findings may not apply to other sites with different practices, different case mix, or 219 different bacterial epidemiology. Third, the management of LVAD-related infections was not 220 standardized during the study period. Indeed, patients were managed on a case-by-case basis, 221 by the local endocarditis team. This could have led to heterogeneity from one site to another, 222 although endocarditis teams follow the same basic principles, have access to similar 223 diagnostic tools and anti-infective treatment, and prioritize evidence-based medicine when 224 225 available.

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#### 227 Conclusions

LVAD is a significant progress that improves survival status, functional capacity, and quality of life, but it carries a high risk of infectious complications, evaluated at 22.6% at one year in this study. With systemic antibacterial treatment for all patients, and debridement in selected cases, the final outcome is usually satisfactory, as most cases of LVAD-related infections may be controlled, with no sequel, and no contraindication for heart transplant.

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#### 235 **Transparency declaration and financial support statement**

236 No external funding was received for this study. All authors: no potential conflicts of interest.

#### 237 Acknowledgments

We are indebted to all the patients who participated in the study, and to the health care workers who took care of them in the departments of cardiology, cardiac surgery, and infectious diseases.

#### References

- Mozaffarian D, Benjamin EJ, Go AS, Arnett DK, Blaha MJ, Cushman M, et al. Heart disease and stroke statistics-2016 update: a report from the American Heart Association. Circulation 2016;133(4):447-54
- 2. Ponikowski P, Voors AA, Anker SD, Bueno H, Cleland JG, Coats AJ, et al. 2016 ESC Guidelines for the diagnosis and treatment of acute and chronic heart failure: The Task Force for the diagnosis and treatment of acute and chronic heart failure of the European Society of Cardiology (ESC). Eur Heart J 2016;37(27):2129-200
- Heidenreich PA, Albert NM, Allen LA, Bluemke DA, Butler J, Fonarow GC, et al. Forecasting the impact of heart failure in the United States: a policy statement from the American Heart Association. Circ Heart Fail 2013;6:606–19.
- 4. Bleumink GS, Knetsch AM, SturkeNboom MC, Straus SM, Hofman A, Deckers JW, Witteman JC, Stricker BH. Quantifying the heart failure epidemic: prevalence, incidence rate, lifetime risk and prognosis of heart failure The Rotterdam Study. Eur Heart J 2004;25:1614-19.
- 5. Agence de la Biomédecine. Rapports annuels d'activité 2015. <u>https://www.agence-biomedecine.fr/Rapports-annuels-d-activite-2015</u>
- 6. Mosterd A, Hoes AW. Clinical epidemiology of heart failure. Heart 2007;93:1137–46.
- Holman WL, Naftel DC, Eckert CE, Kormos RL, Goldstein DJ, Kirklin JK. Durability of left ventricular assist devices: interagency registry for mechanically assisted circulatory support (INTERMACS) 2006-2011. J Thorac Cardiovasc Surg 2013;146:437-41.
- 8. Rose EA, Geligns AC, Mosckowitz AJ, Heitjan DF, Stevenson LW, Dembitsky W, et al. Randomised Evaluation of Mechanical Assistance for the Treatment of Congestive Heart Failure (REMATCH) Study Group. Longterm mechanical left ventricular assistance for end-stage heart failure. N Engl J Med 2001;345:1435-43.

- 9. Wus L, Manning M, Entwistle JWC. Left ventricular assist device driveline infection and the frequency of dressing change in hospitalized patients. Heart Lung 2015;44:225-29.
- 10. Zierer A, Melby SJ, Voeller RK, Guthrie TJ, Ewald GA, Shelton K, et al. Late-onset driveline infections: the Achilles' heel of prolonged left ventricular assist of prolonged left ventricular assist device support. Ann Thorac Surg 2007;84:515-20.
- 11. Xie A, Phan K, Yan TD. Durability of continuous-flow left ventricular assist devices: a systematic review. Ann Cardiothorac Surg 2014;3:547-56.
- 12. Topkara VK, Kondareddy S, Malik F, Wang IW, Mann DL, Ewald GA, Moazami N. Infectious complications in patients with left ventricular assist device: etiology and outcomes in the continuous-flow era. Ann Thorac Surg 2010;90:1270-77.
- 13. Koval CE, Thuita L, Moazami N, Blackstone E. Evolution and impact of driveline infection in a large cohort of continuous-flow ventricular assist device recipients. J Heart Lung Transplant 2014;33:1164-72.

14. Interagency Registry for Mechanically Assisted Circulatory Support (INTERMACS) user's guide. Manual of operation, version 2.3:66. <u>http://www.uab.edu/medicine/intermacs</u>

- 15. Aslam S, Hernandez M, Thornby J, Zeluff B, Darouiche RO. Risk factors and outcomes of fungal ventricular assist-device infections. Clin Infect Dis 2010;50:664-71.
- 16. Kirklin JK, Naftel DC, Pagani FD, Kormos RL, Stevenson LW, Blume ED, et al. Seventh INTERMACS annual report: 15,000 patients and counting. J Heart Lung Transplant 2015;34:1495-04.
- Califano S, Pagani FD, Malani PN. Left ventricular assist device-associated infections. Infect Dis Clin North Am 2012;26:77-87
- 18. Koval CE, Rakita R, AST Infectious Disease Community of Practice. Ventricular assist device related infections and solid organ transplantation. Am J Transplant 2013;13:348-54

- 1 Table 1. Characteristics of patients with left ventricular assist device-related infections
- 2 (n=36) by time of implantation

Variables	
Demographic characteristics	
Age at implantation (years)	$51 \pm 11$
Male	31 (86)
Comorbidity	
Immunocompromised	4 (11)
Diabetes mellitus	4 (11)
Chronic alcoholism	12 (33)
Left ventricular ejection fraction (%)	$22 \pm 7$
Body mass index $(kg/m^2)$	$25.4 \pm 4.9$
Aetiology of heart failure	
Ischemic cardiomyopathy	22 (61)
Dilated cardiomyopathy	14 (39)
Duration of heart failure	
< 12 months	17 (48)
1-5 years	9 (26)
> 5 years	9 (26)
Indication of LVAD implantation <sup>a</sup>	
Bridge-to-transplantation	22 (65)
Bridge-to-recovery	8 (23)
Destination therapy	4 (12)
INTERMACS profile <sup>b</sup>	
I	21 (58)
Ш	0 (0)
	5 (14)
IN IV	3 (8)
V-VII	7 (20)
V-VII	7 (20)
LVAD device	
Heartmate II (Thoratec)	33 (92)
Others <sup>c</sup>	3 (8)

3 Categorical data are summarized as Number (%) of patients and continuous data are

4 summarized as mean  $\pm$  standard deviation.

5 LVAD, left ventricular assist device; INTERMACS, Interagency Registry for Mechanically

6 Assisted Circulatory Support

<sup>&</sup>lt;sup>a</sup> Data available for 34 patients; <sup>b</sup>, INTERMACS profile I = critical cardiogenic shock, II =

<sup>8</sup> progressive decline on inotropic support, III = stable, but inotrope dependent, IV = resting

<sup>9</sup> symptoms, V = exertion intolerant, VI = exertion limited, VII = adavanced New York heart

<sup>10</sup> association class 3; <sup>c</sup> Heartware, Thoratec (n=2); VentrAssist, Ventracor (n=1)

## 1 Table 2

2 Characteristics and management of left ventricular assist device-associated infections (n=36)

Variables	
Clinical manifestation	
Time from implantation to infection, months	2.9 [1.8-7.5]
Purulent drainage	31 (86%)
Pain	17 (47%)
Erythema	22 (61%)
Fever	28 (80%)
Infection site <sup>a</sup>	
Percutaneous driveline	17 (49%)
Locoregional	13 (37%)
Pump	5 (14%)
Laboratory variables	
White Blood Cell count, $10^9/L$	11.8 [5.6-27.7]
Serum C Reactive Protein, mg/l	26 [3-362]
Positive blood culture(s)	4 (10%)
Microbiology	
Staphylococcus aureus	20 (56%) <sup>b</sup>
Enterobacteriaceae	12 (33%) <sup>c</sup>
Pseudomonas aeruginosa	10 (28%)
Coagulase-negative staphylococci	6 (18%)
Corynebacterium sp.	6 (18%)
Others	3 (8%) <sup>d</sup>
Management	
Systemic antibacterial treatment <sup>e</sup>	36 (100%)
Median duration (days)	40 [14-120]
Surgical debridement	17 (47%)

3 Categorical data are summarized as Number (%) of patients and continuous data are

- 4 summarized as median values [first quartile-third quartile].
- <sup>a</sup> data available for 35 patients
- 6 <sup>b</sup> including 2 (10%) methicillin-resistant *Staphylococcus aureus*
- <sup>c</sup> including 7 (58%) extended spectrum beta-lactamase producers

8 <sup>d</sup> Acinetobacter baumanii, Haemophilus aprophilus, and Stenotrophomonas malthophilia,

9 one patient each

<sup>e</sup> Antibacterial treatment included cloxacillin (n=21), pristinamycin (n=15), ciprofloxacin

11 (n=12), gentamicin (n=12), amikacin (n=9), vancomycin (n=7), imipenem (n=7), rifampin

- 12 (n=6), piperacillin-tazobactam (n=5), and ofloxacin, amoxicillin-clavulanate, amoxicillin,
- 13 trimethoprime-sulfamethoxazole, clindamycin, doxycyclin, fusidic acid, ceftazidime,
- 14 ceftriaxone, cefamandole, and linezolid (n<5 for each).