

Safety of Tedizolid as Suppressive Antimicrobial Therapy for Patients With Complex Implant-Associated Bone and Joint Infection due to Multidrug-Resistant Gram-Positive Pathogens: Results From the TediSAT Cohort Study

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A prospective cohort study was conducted to evaluate long-term safety of tedizolid as suppressive antimicrobial treatment in patients with implant-associated bone and joint infection caused by multidrug-resistant gram-positive pathogens. Seventeen patients received tedizolid with a median duration of treatment of 6 months. No patients developed a serious adverse event.

Keywords. bone and joint infection; linezolid; periprosthetic joint infection; suppressive antimicrobial therapy; tedizolid.

Infections are one of the most dramatic complications after an arthroplasty or osteosynthesis following a fracture [1]. These infections are difficult to treat, requiring prolonged treatment courses, with frequent relapses, despite a dedicated medico-surgical management. In cases when complete microbiological eradication is not possible, suppressive antimicrobial therapy

(SAT) may be required, especially if further surgery is not feasible in patients with significant comorbidities or advanced age [2].

Staphylococcus aureus is the cause in the majority of orthopedic-implant infections, but less virulent organisms, such as coagulase-negative *Staphylococcus* (CoNS), are also frequently implicated. The prevalence of multidrug-resistant (MDR) gram-positive organisms, such as methicillin-resistant *S. aureus* and vancomycin-resistant enterococci (VRE), has been steadily increasing, with methicillin-resistant *Staphylococcus epidermidis* identified more frequently through the years [1, 3].

Linezolid (LZD), an oxazolidinone antibiotic, remains active against these MDR gram-positive pathogens in most cases. Nevertheless, the risk of myelotoxicity and polyneuropathy, predominately associated with long-term use, and drug–drug interactions with serotonergic agents are common issues in patients with chronic complex bone and joint infections (BJIs) [4]. Tedizolid (TZD), a newer oxazolidinone agent, has been associated with lower rates of myelotoxicity and drug–drug interactions, and therefore may be a viable alternative [4, 5]. Indeed, TZD is thought to be less active on mitochondrial protein synthesis than LZD, and consequently fewer side effects are expected [6]. Tedizolid has only been validated for the treatment of acute bacterial skin and skin-structure infections, and treatment duration in those approved indications is short (6 days) [7]. To our knowledge, the longest treatment duration with TZD in BJI reported was 12 weeks, with several other case reports of longer treatment durations for variable types of infections [8–10]. The aim of this study was to describe our experience with TZD as SAT in patients with complex implant-associated BJI due to MDR gram-positive pathogens.

METHODS

A prospective monocentric cohort study was conducted between 2017 and 2020 at our referral center for the management of complex BJI (Centre de Référence des Infections Ostéo-Articulaires complexes [CRIOAc]) in Lyon, France (<http://www.crioac-lyon.fr>).

Periprosthetic joint infections and osteosynthesis-associated infections (OAIs) were defined according to the MusculoSkeletal Infection Society 2018 criteria [11] and Fracture-Related Infection Consensus Group [12], respectively. The clinical situation of every adult patient referred to our reference center with PJI or OAI was discussed during multidisciplinary meetings (infectious disease specialist, microbiologist, orthopedic and plastic surgeons). In all cases, use of tedizolid was validated as the last oral treatment option for SAT, due to intolerance, allergy, or resistant pathogen

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to other available oral antibiotics. Notably, the choice of TZD was driven by antimicrobial susceptibility results of the current and previous infectious episodes, if considered as not cured.

SAT was defined as an indefinite administration of antibiotics without curative intention in the context of a chronic infection that would normally require implant removal; the aim of SAT was to prevent infection progression and thus, the loss of limb function. There were no exclusion criteria.

The primary objective of this study was to evaluate the safety of TZD as SAT. Any adverse event (AE), defined as an untoward medical event possibly associated with the use of TZD and graded according to the National Cancer Institute Common Terminology Criteria for Adverse Events, any reason for discontinuation, and standard biological data were prospectively collected. Failure of SAT was defined as the presence of clinical signs suggestive of uncontrolled infection and the need for a new surgical procedure. Anemia was defined as a decrease in hemoglobin level of ≥ 2 g/dL, thrombocytopenia as a platelet count $<150 \times 10^9/L$, leukopenia as a white blood cell count $<4 \times 10^9/L$, and neutropenia as an absolute neutrophil count $<1.5 \times 10^9/L$.

Data were analyzed with Stata 16.1 software (StataCorp, College Station, Texas). Categorical variables were described by counts and percentages, while mean and standard deviation or median and interquartile range (IQR) were used to summarize continuous variables. Biological variables were compared at baseline and 12 months using paired *t* test. Significance was set at $P < .05$.

Patient Consent Statement

This study was subject to declaration with the local Commission for Data Protection and Liberties under the number 20_232 and was recorded on ClinicalTrials.gov under the number NCT04662736.

This study was part of the Lyon BJI cohort study, and patients' consent was obtained for each inclusion.

RESULTS

Between 2017 and 2020, 17 patients received TZD orally at 200 mg once daily (no adjustment for liver or renal function was made) as SAT for late complex BJI, with a median duration of TZD treatment of 6 months (range, 1–31 months; IQR, 2–15 months). Only 1 patient was considered to have failed TZD treatment at 1 month, but 2 others had also received a short course of treatment at the time of data collection, explaining the IQR from 2 to 15 months. Patients were predominantly male ($n = 13$ [76%]), with a median age of 73 years (IQR, 69–81 years), a mean body mass index of 28.1 ± 5.1 kg/m² (range, 19.5–36.7 kg/m²), and a mean American Society of Anesthesiologists score of 2.2 ± 0.6 (range, 1–3). Knee PJIs were the most frequent infections ($n = 10$ [59%]), followed by hip

($n = 5$ [29%]) and shoulder PJI ($n = 1$ [6%]). There was 1 femoral intramedullary nail infection (6%) (Table 1).

Pathogens included MDR CoNS ($n = 16$ [76.2%]), followed by *Corynebacterium striatum* ($n = 2$), vancomycin-resistant *Enterococcus faecium* (VRE; $n = 1$), and/or methicillin-sensitive *S aureus* ($n = 2$). Coinfections with gram-negative pathogens (*Pseudomonas aeruginosa*, *Serratia marcescens*, and *Proteus mirabilis*) were observed in 3 patients. All antimicrobial susceptibilities are found in Supplementary Table 1.

An empirical intravenous (IV) antibiotic therapy followed each surgical procedure (mainly debridement and implant retention; $n = 13$ [76.5%]), with subsequent adjustment based on bacterial susceptibility. Median duration of this primary IV treatment was 47 days (IQR, 35–79 days; range, 5–168 days), followed by LZD in 13 (76.5%) patients. Antibiotic treatment was then changed to tedizolid, including 9 (75.0%) patients who experienced linezolid-induced serious AE: myelotoxicity in 8 (66.7%) patients and severe gastrointestinal intolerance in 1 patient (Table 1). All events were reversible after transition to TDZ. In 4 other patients, TZD was introduced in first intention for SAT after discussion during a multidisciplinary meeting.

The only reason for discontinuation of TZD was failure of the conservative strategy in 4 patients (23.5%), mostly ($n = 3$ [17%]) due to persistence of sinus tract infection. Two patients presented with a small and intermittent sinus tract; it was considered beneficial to continue TZD and they were not considered as failure. One patient presented a new infection of his arthroplasty, with a new gram-negative pathogen (*Citrobacter koseri*). No serious AE or discontinuation of TZD due to an AE was observed. There were no clinically significant drug–drug interactions despite coadministration with serotonergic agents: tricyclic antidepressants ($n = 2$) and tramadol ($n = 4$).

At 12 months after surgery, there was no difference from baseline in platelet count ($P = .55$), white blood cell count ($P = .75$), or neutrophil count ($P = .93$) (Figure 1, Supplementary Table 2). Hemoglobin level increased during the first year of treatment (mean difference of 2.95 ± 3.55 g/dL from baseline), but it was not statistically significant ($P = .051$). Anemia was observed in 2 patients, in whom an alternate etiology was already known (chronic leukemia and esophageal varices). One case of thrombocytopenia, already observed before introduction of TZD in a cirrhotic patient (platelet count of 80 [160 – 400] $\times 10^9/L$), remained stable during a treatment course of 12 months. Transient mild neutropenia ($1.4 \times 10^9/L$, WBC count of 2.1×10^9) was observed in 1 patient. No neurological or gastrointestinal AEs were observed.

Median duration of follow-up was 8 months (IQR, 5–17 months). One patient died during the study period, with death not considered associated with TZD, AE, or uncontrolled infection. Thus, at the end of the study, except for the 4 patients considered as having failed treatment and the 1 patient who died, TZD was continued in all remaining patients with control of their infection.

Table 1. Patient Demographics and Outcomes of Late Chronic Infections Under Suppressive Antimicrobial Therapy With Tedizolid

Patient ID	Sex	Age at Infection, y	BMI, kg/m ²	ASA Score	Concomitant Drug Administration ^a	Site of Infection	Type of Surgery	Pathogens	LZD AE	Days of LZD Prior to Switch to TZD	Duration of TZD, mo (d)	Reason for Stopping TZD	TZD AE	Follow-up duration, mo (d)	Outcome
1	M	71	27.8	1	Amitriptyline, tramadol	Knee PJI	DAIR	MRSE	NA	NA	31 (960)	NA	None	38 (1168)	Infection controlled
2	M	86	22.0	3	None	Knee PJI	No surgery (inextirpable implant)	History of various pathogens infection (<i>Helicobacterium</i> , <i>Enterococcus faecalis</i> , MSSA, <i>Corynebacterium striatum</i> , <i>Proteus mirabilis</i> , <i>Escherichia coli</i>)	NA	NA	6 (210)	Persistent discharge, uncontrolled infection. TZD stopped as a consequence of a serious AE (seizure) due to ertapenem (concomitant administration)	None	9 (281)	Failure
3	M	79	26.8	2	Tramadol	Knee PJI	DAIR	No growth (history of MRSE infection)	Diarrhea Asthenia	29	16 (516)	Persistent discharge, pain, positive culture despite SAT	None	17 (545)	Failure
4	M	82	28.9	3	Tramadol	Knee PJI	DAIR	MRSE	Thrombopenia (63 x 10 ⁹ /L)	19	17 (540)	NA	None	18 (559)	Infection controlled
5	M	74	26.6	2	None	Hip PJI	DAIR with ceramic head exchange	MRSE	Anemia (6.9 g/dL)	28	2 (61)	New PJI	None	5 (153)	Failure
6	M	64	29.4	2	None	Knee PJI	DAIR	MSSE <i>Staphylococcus hominis</i> (MS) (history of MRSE infection)	NA	NA	14 (441)	NA	None	17 (524)	Infection controlled
7	M	71	28.4	2	None	Knee PJI	Arthrodesis	MRSE	None	28	6 (190)	NA	None	7 (243)	Infection controlled
8	M	69	32.0	2	None	Hip PJI	DAIR	MRSE <i>Pseudomonas aeruginosa</i>	None	35	5 (175)	NA	None	8 (256)	Infection controlled
9	F	89	20.3	2	None	Knee PJI	DAIR	MRSE	Neutropenia	14	21 (645)	NA	None	21 (659)	Small fistula ^b
10	F	87	34.2	3	None	Knee PJI	DAIR	MRSE	Thrombopenia (16 x 10 ⁹ /L)	24	8 (244)	NA	None	8 (269)	Small fistula ^b
11	M	81	36.1	3	None	Shoulder PJI	DAIR	MSSA (history of MRSE infection)	Anemia (8.6 g/dL) Thrombopenia (103 x 10 ⁹ /L)	5	15 (483)	NA	None	15 (488)	Infection controlled
12	F	24	19.5	2	None	Femoral intramedullary nail	Nail exchange	<i>Staphylococcus capitis</i> <i>Enterococcus faecium</i> (VR)	Anemia (6.9 g/dL)	56	13 (423)	NA	None	16 (498)	Infection controlled

Table 1. Continued

Patient ID	Sex	Age at Infection, y	BMI, kg/m ²	ASA Score	Concomitant Drug Administration ^a	Site of Infection	Type of Surgery	Pathogens	LZD AE	Days of LZD Prior to Switch to TZD	Duration of TZD, mo (d)	Reason for Stopping TZD	TZD AE	Follow-up duration, mo (d)	Outcome
13	F	70	28.4	2	Amitriptyline	Hip PJI	DAIR	MSSE <i>C striatum</i> (PCR)	Anemia (8.1 g/dL) Thrombopenia (131 x 10 ⁹ /L)	20	6 (212)	NA	None	7 (236)	Infection controlled
14	M	67	22.5	2	None	Hip PJI	1-stage exchange	MRSE MSSA <i>P mirabilis</i>	Thrombopenia (75 x 10 ⁹ /L)	28	2 (87)	NA	None	5 (157)	Infection controlled
15	M	77	31.1	2	None	Knee PJI	DAIR	<i>Staphylococcus caprae</i> (MR) (history of MSSA, <i>S capitis</i> [MR] infection)	None	14	1 (35)	Persistent discharge, uncontrolled infection	None	2 (78)	Failure
16	M	54	36.7	2	None	Hip PJI	DAIR	MRSE (history of MRSA)	None	14	1 (36)	NA	None	3 (93)	Infection controlled
17	M	73	27.1	3	Tramadol	Knee PJI	DAIR	<i>C striatum</i>	NA	NA	1 (50)	NA	None	3 (97)	Infection controlled

Abbreviations: AE, adverse event; ASA, American Society of Anesthesiologists; BMI, body mass index; DAIR, debridement, antibiotic, implant retention; F, female; ID, identifier; LZD, linezolid; M, male; MR, methicillin-resistant; MRSA, methicillin-resistant *Staphylococcus aureus*; MRSE, methicillin-resistant *Staphylococcus epidermidis*; MS, methicillin-susceptible; MSSA, methicillin-susceptible *Staphylococcus aureus*; MSSE, methicillin-susceptible *Staphylococcus epidermidis*; NA, not applicable; PCR, polymerase chain reaction; PJI, periprosthetic joint infection; SAT, suppressive antimicrobial therapy; TZD, tedizolid; VR, vancomycin-resistant.

^aOnly considered drugs with potential drug–drug interactions with oxazolidinones: tricyclic antidepressant, selective serotonin reuptake inhibitors, and tramadol. No other drugs with potential interaction were prescribed to patients.

^bThese 2 patients were not considered as having failed treatment (see text).

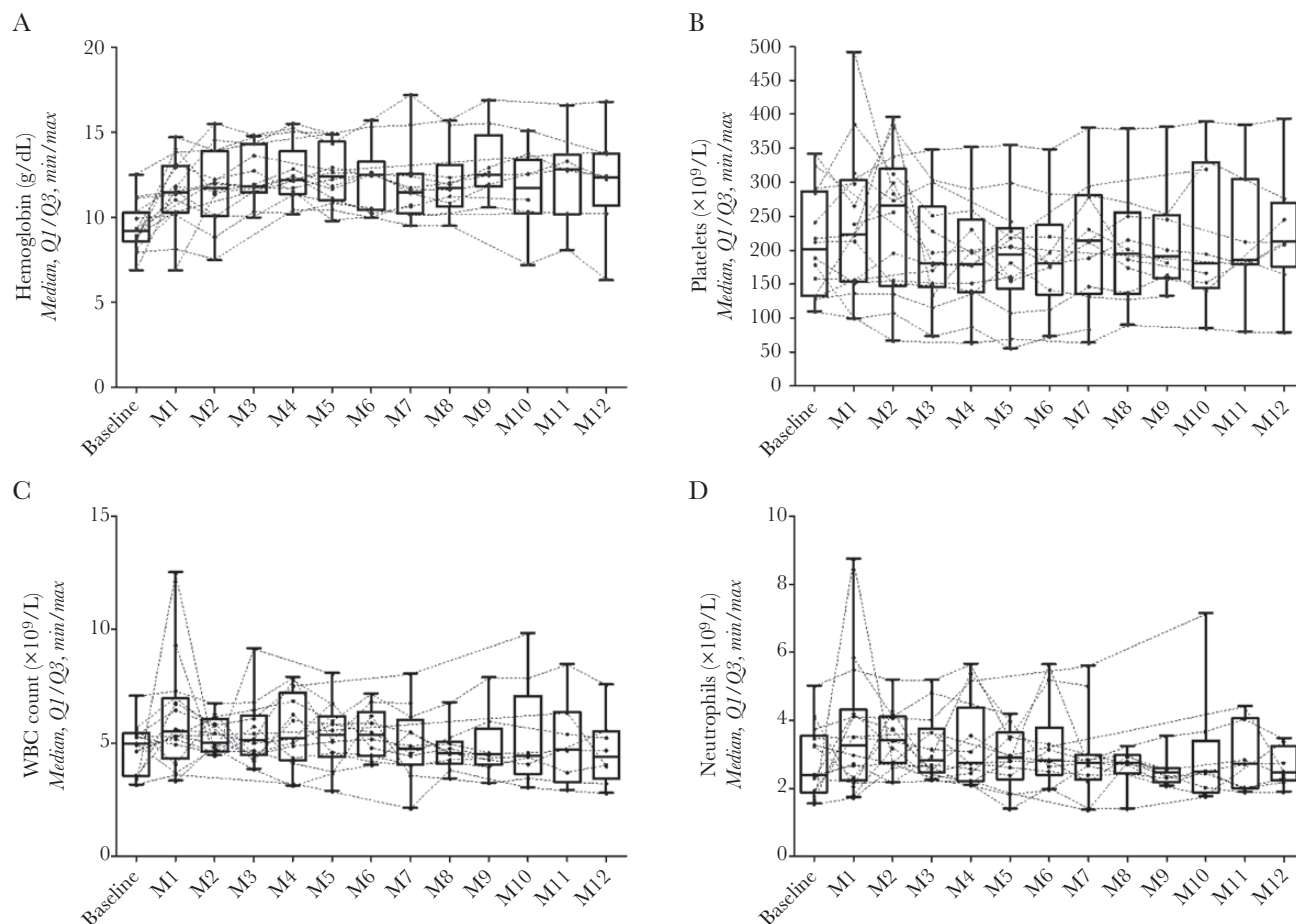


Figure 1. Evolution of hemoglobin (A), platelet count (B), white blood cell (WBC) count (C), and neutrophil count (D) during the first 12 months of suppressive antimicrobial therapy with tedizolid.

DISCUSSION

To our knowledge, this is the longest cohort study of patients who received TZD as an SAT for complex orthopedic implant-associated infections. In this study, TZD was well-tolerated, without any significant hematology- and neurology-related AE, despite the fact that it was initiated because of intolerance to LZD in 9 patients. This suggests that the safety profile of TZD may be more favorable than that of LZD during prolonged treatment courses.

In a clinical trial assessing noninferiority of TZD vs LZD in acute skin and soft tissue infections, a lower incidence of myelotoxicity was observed at the end of treatment with TZD (4.9%) when compared to LZD (10.8%; $P = .0003$) [13]. Furthermore, several case reports and case series have suggested the safety of TZD treatment in patients with LZD-associated myelotoxicity, with resolution of hematologic troubles following the switch to TZD [5].

Recently, 2 other studies on osteoarticular infection treatment by prolonged courses of TZD have been published [8, 9]. In a study on PJI treatment with TZD, patients were treated

with a mean duration of TZD of 8.0 ± 3.27 weeks (range, 6–12 weeks). More than 80% of the patients completed their treatment, and only a few experienced hematological complications, with none leading to discontinuation of TZD [8]. As in our study, their results showed an increase in hemoglobin values, which appears to be related to a restoration process after blood depletion secondary to surgical procedures.

In our cohort study, clinical failure of SAT occurred in 4 patients (23.5%), a rate quite similar to previous studies [2]. It is arguable that 2 patients presenting sinus tract were not considered to have failed treatment. Given the intermittent nature of these small fistula, however, treatment with TZD was considered beneficial to control the infection and to avoid another surgery.

The main limitation of this study is the small size of the population, which precludes definitive safety conclusions, especially regarding rare AEs. Yet, no patient developed AEs leading to treatment discontinuation despite the long duration of the follow-up, and no clinically significant drug–drug interaction was reported. Further clinical investigation is required to confirm the efficacy and safety of TZD as a therapy for BJI.

TZD treatment remains expensive (currently approximately €200 per tablet in France [approximately US\$400]), compared with LZD, which is available as a generic drug and costs approximately €14 per day in France (approximately US\$5). This could be a barrier for TDZ administration as SAT, particularly when considering long treatment courses. In our study, TZD was not provided by the company; in France, this drug is attributed to and delivered by hospitals. However, TDZ is not available in all countries.

It is possible that chronic low-grade implant-associated infections may not always require the administration of long-term suppressive antibiotic treatment. However, considering the high risk of infection progression and devastating consequences on the function of the limb and the patient's life, we believe that SAT is indicated after conservative surgical procedures in those patients without curative alternatives. In addition to TZD, dalbavancin remains an alternative choice in these cases of MDR gram-positive infections. However, this treatment requires IV administration every 1–2 weeks [14], and it has been decided not to use this option at our center.

In conclusion, TZD seems to be a safe option as SAT in patients with complex implant-associated BJI due to MDR gram-positive pathogens, if no other safe oral antibiotic treatment option is available.

Supplementary Data

Supplementary materials are available at *Open Forum Infectious Diseases* online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

Notes

Potential conflicts of interest. T. F. received speaker's fees from MSD in 2021. All other authors report no potential conflicts of interest.

All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

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