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Review

Oral Flucloxacillin for Treating Osteomyelitis: A Narrative Review of Clinical Practice

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Abstract

Flucloxacillin (FLU) administered by the oral route is widely used for treating various infections, but there are no published retrospective or prospective trials of its efficacy, or its advantages or disadvantages compared to parenteral treatment or other antibiotics for treating osteomyelitis. Based on published in vitro data and expert opinions, other non-β-lactam oral antibiotics that have better bone penetration are generally preferred over oral FLU. We reviewed the literature for studies of oral FLU as therapy of osteomyelitis (OM), stratified by acute versus chronic and pediatric versus adult cases. In striking contrast to the prevailing opinions and the few descriptive data available, we found that treatment of OM with oral FLU does not appear to be associated with more clinical failures compared to other oral antibiotic agents. Because of its narrow antibiotic spectrum, infrequent severe adverse effects, and low cost, oral FLU is widely used in clinical practice. We therefore call for investigators to conduct prospective trials investigating the effectiveness and potential advantages of oral FLU for treating OM.

Key words: oral flucloxacillin; osteomyelitis; literature review; clinical treatment; clinical remission

Introduction

The traditional antibiotic regimen for a patient who has undergone surgical debridement for osteomyelitis (OM) includes 2-6 weeks of parenterally administered agent(s), followed by an oral course of several weeks (1,2). Parenteral (usually intravenous) antibiotic therapy is associated with several substantial problems: it requires either prolonged hospitalization or treatment by an outpatient parenteral antibiotic therapy (OPAT) unit; the required intravenous catheter (usually a central line) is associated with frequent adverse effects (including infection and thrombosis); and, it is associated with high financial costs. For example, an evaluation by Gardiol et al. (3) of their OPAT program in Switzerland found that 16 of their 179 (9 %) OPAT patients had complications associated with the treatment. The most common were drug-related adverse events, which occurred in ten patients (5.5 %) and required readmission to the hospital in three cases. There were also six line-related (all peripheral) adverse events (3.5 %) (3).

Flucloxacillin (FLU) is a narrow-spectrum,
semisynthetic penicillin available in both parenteral and oral formulations (4-8). It is active against aerobic gram-positive pathogens, notably including methicillin-susceptible *Staphylococcus aureus* (MSSA) (8). FLU is closely related to other semisynthetic penicillins (nafcillin, oxacillin, (di)cloxacillin) that have been widely used for decades for a variety of orthopedic and community-acquired skin infections worldwide (8,9). Treatment with FLU has several theoretical advantages over other antibiotic agents for these infections: it has bactericidal properties (potentially important for selected infections); its antibiotic spectrum is limited (reducing risk of antibiotic resistance); it is inexpensive (reducing financial costs); and the two formulations afford the possibility of an early switch from intravenous to oral dosing (reducing risks and potentially allowing earlier hospital discharge) (6). One potentially challenging issue is that the bone penetration of oral β-lactam antibiotics, at least in *vitro*, is poor (4-7). Unfortunately, there are almost no published clinical data on the effectiveness or safety of treatment with oral FLU for osteomyelitis (OM).

This lack of evidence of the usefulness of oral semi-synthetic antibiotic agents has led to international expert groups and guidelines, such as the Infectious Diseases Society of America (IDSA) guidelines (9) and the European Society for Pediatric Infectious Diseases (ESPID) (10) to recommend that the preferred route of administration for treating OM are intravenous FLU or a first-generation cephalosporin (9,10). Nevertheless, clinicians in some countries with a low endemicity of methicillin-resistant *S. aureus* isolates, e.g., the Nordic countries, Baltic States and Netherlands, use oral FLU, at least for the maintenance therapy, of staphylococcal OM (6). Several recent studies have demonstrated that oral β-lactam antibiotics, e.g. co-amoxiclav, can be safely used in many types of bone and joint infections, including for diabetic foot OM (11). In the United Kingdom, the prescribing of oral FLU has increased by 21% during the past decade (12), and oral FLU can be used for OM (13). Considering the potential benefits, and lack of published data, we present this narrative review to address the potential place of oral FLU for treating OM.

**Methods**

We performed a narrative literature review by searching for articles in PubMed (using the MeSH terms “oral flucloxacillin”, “*Staphylococcus aureus*” and “osteomyelitis”), Google Scholar and references cited in retrieved articles. We reviewed articles written in English; with correspondence in German, French, Romanian and Turkish languages. Our review only sought papers concerned with oral FLU for treating MSSA infections, not those on intravenous formulations or for treatment of other bacteria. For our inclusion criteria we defined that the duration of oral FLU used in the study must be more than the half of the total antibiotic course. We did not consider antibiotic dosing of oral FLU considerations or minimal clinical follow-up time of enrolled patients as inclusion criteria. We excluded studies: on animals; using combination therapies with oral rifampin; reporting on infected implants or soft tissue infections; on virulence analyses (14); on *S. aureus* body carriage or colonization (15); on pharmacokinetic-pharmacodynamics (PK/PD) issues (16); or, on minimal inhibitory concentration (MIC) assessments (17,18). We also excluded papers predominatingly related to only related antibiotic regimens, such as nafcillin, oxacillin, or (di)-cloxacillin. Additionally, we explored data on the levels of oral FLU use in Switzerland during the past decade.

**Results**

**General results of the literature search**

Our review produced 420 articles, only 58 of which met our criteria and we used these to synthesize our review. Papers with own clinical cases of oral FLU use in OM are analyzed more in detail (Figure 1). We stratified them as involving childhood or adult OM cases, acute or chronic OM cases, and between case reports and case series (Tables 1-5). Although many of the included papers provided information on the feasibility of oral FLU therapy for OM, this was not the underlying primary issue of any. As the heterogeneity and the large amount of missing information in the papers were so great, we could not attempt any type of meta-analyses of our results. Of note, most of the studies were conducted in English speaking (particularly the United Kingdom) and Nordic countries.

**Authority recommendations for prescription of oral flucloxacillin in Switzerland**

According to the most frequently used Swiss Federal documentation (40), general indications for treatment with FLU are the same as in most other countries: skin and soft tissue infections, infection with MSSA, susceptible coagulase-negative staphylococci, as well as streptococci and other non-staphylococcal gram-positive bacteria (*albeit as a weak recommendation*). Importantly, they do not specify if the route of administration routes should be different for bone than for soft tissue infections. Among Swiss Infectious Diseases physicians, treating
with oral FLU for OM is unpopular for the treatment of OM (personal communications). One example of a related consequence is that in Switzerland, because prospective trials involving use of oral antibiotic therapy must pass through the Ethical Committees, researchers avoid using oral FLU as a therapeutic option for studies of bone and joint infections (41). This is not the same in other countries (42). For example, in Sweden, Fiji and Australia, oral FLU can be officially used for studies of treatment of OM (43-46).

**Official prices and registered use of oral flucloxacillin in Switzerland**

The daily price for treatment with oral FLU (500 mg, 3-4 times a day) on the Swiss market is five Swiss Francs ($5 US dollars in October 2019). The quantity of oral FLU prescribed in Switzerland between 2004 and 2017 was 0.13 DDD (defined daily dose) per 1000 patient-months. There was a two-fold increase in use during this time period in the French-speaking part of the country, a 29% increase in the Italian-speaking part, and an 18% in the German-speaking majority regions (ANRESIS databases; 47). In our Balgrist University Hospital, a referral center for orthopedic infections, the corresponding DDD was 0.72 per 1000 patient-months in 2017. In comparison, in the United Kingdom (the country with the most publications regarding oral FLU in OM), the average monthly prescription rates increased from 4.74 per 1000 patient-months in 2004 to 5.74 in 2013, an increase of 21.1% (12). We do not know the reasons for this increase in prescribing of oral FLU, but it might be that British physicians chose to prescribe more of the 500 mg tablets the 250 mg capsules (12). In Sweden, the usual daily dosing of oral FLU is 1.5 g three times daily for OM and septic arthritis (44). The EUCAST Clinical Breakpoint Tables (European dosing recommendations) promote a “standard dose” of oral FLU of 1.0 g three times daily; and a “high oral dose” of 1g four times daily in some cases (48).

**International guidelines**

To our knowledge, treatment with oral FLU is not a primary choice for OM in any internationally accepted guidelines. For example, the ESPID guidelines (10) recommend initial therapy with a 1st or 2nd generation cephalosporin (for other susceptible bacteria other than MSSA). For treating adult in the OM population, the IDSA guidelines (9) advocate 1.5-2.0 g of intravenous oxacillin (the nearest equivalent to flucloxacillin) every four hours. Both guidelines consider almost all other oral β-lactams, as a group and independently of their dosing, inappropriate due to poor bone penetration in vitro (with the possible exception of amoxicillin and co-amoxiclav in certain circumstances); this is true even when the patient has shown a good response to initial parenteral therapy treatment with a non-preferred agent. In striking contrast, after a favorable clinical response to parenteral treatment, guidelines encourage switching to oral antibiotic classes for almost all other agents (e.g., quinolones, linezolid, clindamycin, co-trimoxazole, tetracyclines,
and metronidazole) provided there is (9). The failure to recommend oral FLU also applies for DFO in the US guidelines (49).

Regional and national recommendations

In contrast to their international counterparts, many national guidelines explicitly recommend treatment with oral FLU for MSSA OM, generally citing the favorable personal experience of the author groups (12). A UK study reported that general practitioners treated 394 OM episodes at least partially with oral FLU, with generally favorable results (12). Recent recommendations from South Korea also mention the possibility of post-surgical treatment of OM with oral FLU (50). Swedish (44), Australian (46) and Fijian (45) guidelines also support treatment with oral FLU in OM, including spondylodiscitis (44). Similarly, the national Australian Antibiotic Guidelines (Therapeutic Guidelines) promote oral FLU as a first line treatment for acute and chronic OM, albeit following a 2-4 week course of intravenous FLU (46).

Published clinical efficacy of oral flucloxacillin in osteomyelitis

Tables 1-5 present summaries of the published clinical literature on treatment of OM with oral FLU. Over twenty years ago Lowden et al. reported that among five children with clavicular OM (19) four (80%) relapsed after three weeks intravenous FLU followed by three to four weeks of oral FLU. The author’s attributed this high recurrence rate to the clavicle being a difficult anatomic site to treat. No subsequent studies have reported such a high failure rate, suggesting there may have been other problems in the management of these patients (19). Vinod et al. reported their experience with children treated for OM or septic arthritis (20). In 77% of the cases the children received intravenous FLU followed by oral FLU. Overall, the recurrence rate was only 1.4%, demonstrating the efficacy of oral FLU in treating hematogenous OM (20). Similarly, Nunn et al. demonstrated efficacy in the majority (exact numbers are not provided) of 43 evaluable children treated in an African hospital for OM with oral FLU following initial intravenous therapy (51). In 2001 Beronius et al. published their experience in Sweden of treating 42 episodes of spondylodiscitis for a median of 10 days with intravenous antibiotics (cefuroxime in two-thirds of cases), followed by a median of 179 days of oral FLU, none of which were associated with a relapse (23). For the subset of patients with DFO, we found no well-designed trials comparing oral FLU with other antibiotic regimens. One study specifically comparing patients treated with oral β-lactam antibiotics to those treated with oral quinolones for DFO found no significant difference in the remission rates (52). Several case series, reports and surveys have demonstrated the feasibility of treatment with oral β-lactam therapy in DFO (3,36). Interestingly, a survey of Australasian infectious diseases clinicians found that oral FLU is the maintenance antibiotic treatment of choice, associated with good clinical responses, for staphylococcal DFO (53).

In sum, the experience of the majority publications reporting on outcomes of treatment with oral FLU for OM have demonstrated good efficacy. While these studies used different dosing regimens and treatment durations, and had non-uniform rates of surgical treatment (Tables 1-5), many used oral FLU as a maintenance therapy after a course of initial parenteral antibiotic therapy. Among the reported cases series there were relatively few documented failures (Tables 1, 2 and 4). In one case, the authors treated a pediatric spondylodiscitis with oral FLU (27) and incriminate the relatively short duration of therapy (24 days), not the choice of FLU, as the cause of relapse. Other reported recurrences occurred in studies from New Zealand (31) and Denmark (36).

Drug interactions and tolerability of oral flucloxacillin in the treatment of osteomyelitis

Although comparative large epidemiological data are lacking, according to personal experience of various authors (personal communications) and published literature, oral FLU probably causes not more adverse events than other penicillins. The most common reported side effects include hypersensitivity, gastrointestinal symptoms, exanthema, nephrotoxicity, myelotoxicity, oxaproline acidosis. Drug-drug interactions have been reported with paracetamol (acetaminophen) (40,42), and rarely with warfarin is (54) and rifampin. In a study in 15 adult patients with various osteoarticular infections (55) the combined treatment with intravenous FLU and oral rifampicin significantly increased serum FLU levels by 45% in 10 patients, while a decrease was observed in one patient. The clinical significance of this finding remained unknown, since all infections were cured (55). It is likely that oral FLU could also be associated with such an interaction, although possibly at a subclinical level. Contrary to common belief, it is probably not necessary to administer oral FLU on an empty stomach, as concomitant food intake does not appear to alter the pharmacokinetic parameters (56). One other interaction of note is that combining treated with FLU (like other penicillins) with probenecid commonly can boost the serum FLU concentrations (8,40,42).
Table 1 - Case series in humans of chronic osteomyelitis

| Reference, Year (Country) | Patients, Age | IV treatment | Duration of oral FLU | Follow-up | Outcome |
|---------------------------|---------------|--------------|---------------------|-----------|---------|
| Lowden et al, 1997 (New Zealand) | 5 patients | 60% drainage | 21-42 days | 0.5-1 year | 80% relapse recovery not documented |
| Vinod et al, 2001 (United Kingdom) | 32 patients; 3 months to 16 years | 49% drainage | 21-42 days | 66% > 12 months | relapse rate 1.4% |
| Sur et al, 2015 (United Kingdom) | 55 patients | 30% debridement | FLU IV, vancomycin | not known | not known |
| Torda et al, 2019 (Malawi) | 3 patients | not known | 6 weeks | not known | no relapse |
| Beranius et al, 2001 (Sweden) | 42 patients | median 10 days | 179 days | not known | no relapse |

FLU = Flucloxacillin; IV = intravenous; MSSA = Methicillin-susceptible Staphylococcus aureus.

Table 2 - Case reports of treatment of children with acute osteomyelitis

| Reference, Year (Country) | Pathogenesis | Pathogen | Treatment Surgery, IV antibiotic | Duration of Oral FLU | Outcome |
|---------------------------|--------------|----------|----------------------------------|---------------------|---------|
| Hughes et al, 2016 (United Kingdom) | Hematogenous | MSSA | No operation | 14 days | no relapse |
| Pabla et al, 2011 (Malawi) | Traumatic | Unknown | No operation | 64 days | no relapse |
| Chiappini et al, 2016 (Italy) | Microtrauma | MSSA | Drainage | 28 days | no relapse |
| Dahal et al, 2019 (Nepal) | Spondylodiscitis | MSSA | 10 days FLU | 14 days | 1 relapse |

FLU = Flucloxacillin; IV = intravenous; MSSA = Methicillin-susceptible Staphylococcus aureus.

Table 3 - Case reports of treatment of children with chronic osteomyelitis

| Study | Pathogenesis | Pathogen | Treatment Surgery, IV antibiotic | Duration of oral FLU | Outcome |
|-------|--------------|----------|----------------------------------|---------------------|---------|
| El Mezouar et al, 2014 (Morocco) | Trauma | MSSA | No operation | 120 days | no relapse |
| Freeman et al, 2008 (Malawi) | Trauma | No culture | not known | 12 days FLU | no relapse |
| Rajakulendran et al, 2013 (United Kingdom) | PIN-tract infection | MSSA | Not known | 85 days | no relapse |
| Kerr et al, 2019 (United Kingdom) | Acetabular osteomyelitis | MSSA | Not known | 28 days | no relapse |

FLU = Flucloxacillin; IV = intravenous; MSSA = Methicillin-susceptible Staphylococcus aureus.

Table 4 - Case reports of treatment of adults with acute osteomyelitis

| Study | Pathogenesis | Pathogen | IV treatment | Duration of oral FLU | Recovery |
|-------|--------------|----------|--------------|---------------------|---------|
| Waterhouse et al, 2013 (New Zealand) | Hematogenous | MSSA, H. parainfluenzae | 6 days Augmentin PO 2 days FLU IV | not known, long | 1 relapse |
| Cheer et al, 2009 (UK) | Microtrauma | MSSA | 6 weeks FLU | not known, long | No relapse |
| Marshman et al, 2008 (UK) | Hematogenous | MSSA | 12 days FLU | 28 days | No relapse |
| Erturan et al, 2012 (UK) | Hematogenous | C. pseudodiphteriticum | Synovectomy, 8 days FLU IV 8 days vancomycin IV | 42 days | No relapse |

C. = Corynebacterium; FLU = Flucloxacillin; IV = intravenous; PO = orally; MSSA = Methicillin-susceptible Staphylococcus aureus.

Table 5 - Case reports of treatment of adults with chronic osteomyelitis

| Study | Pathogenesis | Pathogen | IV treatment | Duration of oral FLU | Recovery |
|-------|--------------|----------|--------------|---------------------|---------|
| Opara et al, 2007 (United Kingdom) | Hematogenous | MSSA, TB | 2 weeks FLU & Fusidic acid IV | 14 days | Residual handicap, no relapse |
| Jeppesen et al, 2015 (Denmark) | Hematogenous | MSSA | No surgery | 84 days | No relapse |
| Lindau et al, 2015 (United Kingdom) | Traumatic | Not known | Gentamicin beads | 56 days | No relapse |
| Torda et al, 1995 (Australia) | Spondylodiscitis | MSSA | Not known | 280 days | No relapse |
| Padala et al, 2013 (UK) | Spondylodiscitis | MSSA | Fusidic acid | 90 days | No relapse |

FLU = Flucloxacillin; IV = intravenous; MSSA = Methicillin-susceptible Staphylococcus aureus; TB = Tuberculosis

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Discussion

Our review of the literature revealed that despite its widespread use around the world, there is remarkably little documentation of the clinical experience of treatment with oral FLU alone for the treatment of acute or chronic, implant-free, OM in children or adults. We found no randomized trials or retrospective multivariate analyses that adjusted for the large case-mix of included subjects.

The few published reports of case series clearly reflect the personal experience of the authors, and are almost exclusively composed of case reports and small case series. The information on efficacy and adverse events if present at all, is largely hidden within the reports, as the primary objectives of the studies differed from those of our review.

Based on in vitro data, treatment with oral FLU (as with all oral β-lactam agents) warrants careful observation when used for treating OM. In most studies the ratios of bone to intravenous concentrations for β-lactams are about 5-20%, with a very wide inter-individual variation in the availability of oral FLU. As an example, the serum trough levels reported by Gath et al. were 15±5.9 mg/L (5), while the peak levels reported by Dijkmans et al were 22-26 mg/L (range, 7-53 mg/L) (6). Thus, while most intravenous (and probably intramuscular) β-lactams reach the necessary MIC for MSSA in bone (57-61), this might not be the case with oral administration. One cause for the relatively low levels achieved with oral FLU is likely the innate variabilities in its gastrointestinal absorption (6). Among humans, 9% largely fail to absorb FLU and another 17% are poor absorbers (61). A pharmacologic in vivo study from the Netherlands (61) compared levels of cefuroxime and FLU when given by oral versus parenteral routes in plasma and intraoperative bone during twenty hip and knee arthroplasty surgeries. The authors allocated patients to four groups: either 1 x 500 mg or 7 x 500 mg oral cefuroxime, following 2000 mg intravenous FLU; and either 1 x 500 mg or 7 x 500 mg oral FLU followed by 1500 mg parenteral cefuroxime. All oral administrations failed to achieve a measurable osseous concentration (60). FLU is a highly protein bound drug and it is predominantly the free drug that is active in the killing of bacteria. Even with a relatively high dosage of 1 g q.i.d, free plasma drug concentrations might drop below the MIC of a typical strain of MSSA (0.25 mg/l) at 50% of the dosing interval (56). Given a drug penetration into bone ratio of 10-20%, it is unlikely that a given dose of oral FLU will achieve therapeutic concentrations in the bone for several hours. We only found one published study reporting the contrary: Köndell et al. found a mandibular bone concentration of 3.8 mg/L after oral administration of FLU, which would likely be therapeutically adequate for treating MSSA OM (62).

A key issue is that serum antibiotic concentration measurements do not reflect the entire pharmacologic picture, as it is not a highly predictable surrogate for the actual concentration of the administered antibiotic in infected bone (16). Likewise, information is lacking regarding the relationship between serum antibiotic concentrations over time and antibiotic effectiveness (time-over minimal inhibitory concentration [MIC]) for FLU and MSSA, as FLU is not an antibiotic that is peak concentration-dependent (as is true of all β-lactam antibiotics agents). Interested readers can review a paper in which members of our group recently summarized insights on the pharmacokinetic-pharmacodynamics of antibiotics in the bone (16). The effectiveness of antibiotic treatment for OM is complex, depending on the interaction of drug, host, microbial agent and other factors. For example, it is disturbing that Alvarez-Ferrero et al. (61) failed to detect a measurable osseous concentration in vivo after administration of oral FLU, but they did not address the large technical and analytical problems in assessing drug concentrations in bone (61). There are likely also differences in antibiotic concentrations in cancellous versus cortical versus mixed homogenized bone, in infected versus non-infected bone, and issues related to blood contamination during bone sampling. Therapeutic drug monitoring (TDM) might be a solution to concerns about therapeutic concentration. In patients being treated with FLU use (oral or IV), several centers have introduced such TDM policies, but its clinical benefit remains unproven. Most TDM publications to date provide only laboratory (not clinical) data for FLU (63). This contrasts to antibiotics such as cefepime, for which TDM has been shown to help prevent potentially severe adverse events (64).

The outcome of oral FLU treatment of OM does not appear to be worse than with other oral drugs (2), although direct comparative data are lacking. So, why does treatment with oral FLU seem to be effective for OM if its bone penetration appears to be limited? In our review (Tables 1-5), we found few reported failures of OM treatment, either in adults or children. Likewise, in the study by Vinod et al., of 32 children with OM treated with IV FLU for a median of three days, followed by oral therapy for a median of 6 weeks, the relapse rate was only 1.4% (20). We can only speculate on the reasons. Based on our review and personal communications, even in “oral FLU-supporting countries” (44-46), treatment of acute as well as chronic OM is almost always started with
intravenous FLU for about two weeks, which is then followed by a longer period of oral FLU. Official guidelines in Australia (43,46) have been recommending a much longer intravenous duration for OM compared to what is currently in place in Switzerland (~2 weeks or less). It is possible that long oral maintenance antibiotic therapy is unnecessary, as an unimpaired immune system is able to eradicate the remaining bacteria (1,41), including for DFO (44,45,52). This is why in this review we elected to circumvent the confounding bias of long duration intravenous therapy by targeted OM episodes in which ≥50% of the course was with oral FLU.

There is more data supporting the clinical efficacy in treating OM in various settings with oral β-lactams other than FLU. For example, many experts in the Nordic countries consider cloxacillin (which is not FLU, sensu strictu) the preferred oral β-lactam drug. For example, Bachur et al. found good outcomes with a four-day intravenous administration of cefazolin with an early switch to oral cephalaxin for 28 days in stable OM (65). Similarly, Le Saux found no superiority for a long-duration intravenous β-lactam administration in children with hematogenous OM (66) compared to a short duration. Likewise, Zaoutis et al. in a retrospective study of hematogenous pediatric OM, treatment with prolonged intravenous β-lactam therapy compared to an early switch to oral therapy gave similar outcomes at six months (67). Several pediatric studies in the 1970s and 1980s summarized the feasibility of oral β-lactam care for OM (11). The recently published OVIVA trial prospectively randomized various types of severe bone and joint infections, including infected arthroplasties and osteomyelitis, into either six weeks of intravenous antibiotic therapy or a switch to oral antimicrobials after up to 7-days of initial parenteral therapy (7). Roughly 38% of all participants had infections with MSSA. Overall, 74 patients in the oral arm were treated with oral penicillins and clinical failure was noted in 17 (23%), compared to 9 of 57 (16%) in the intravenous arm, a non-significant difference (7; Supplementary Appendix).

There is also a great variability in antibiotics used in patients with DFO, with co-amoxiclav among the most frequently orally administered β-lactams worldwide (11). In a retrospective cohort analysis of a clinical pathway for patients with DFO, the authors conducted a cluster-controlled Cox regression model with an emphasis on the more than 300 patients treated with oral co-amoxiclav (at a median dose of 2000 mg BID/day) (11). After a median follow-up of three years, DFO recurred in 22% of the patients, which is compatible with rates reported for various treatments in the literature. In multivariate analysis, treatment with oral co-amoxiclav did not influence the likelihood of remission or recurrence, either when used as the only agent from the start, or when administered for >50% of the entire antibiotic course (11). Likewise, in a prospective trial by Lázaro-Martínez et al. randomizing patients with DFO to either primarily antibiotic treatment (24 cases with oral co-amoxiclav and 24 cases with other oral antimicrobials) for ninety days or to “conservative” surgery and antibiotics for 10 days (68), there were no significant differences in outcomes. In contrast to the situation with oral FLU, some international guidelines (49) suggest co-amoxiclav is a viable oral antibiotic option for treating DFO (11). These data from studies of other oral β-lactam agents also support the use of oral FLU as a possible alternative in OM.

Our narrative review has several limitations, the most important of which is the heterogeneity of practice regarding treatment with oral FLU. Considering that in some countries oral FLU use is already firmly established and recommended in national guidelines (44-46), there are probably tens of thousands of OM cases treated each year in this way. Unfortunately, these clinicians have rarely published their experience. For this review we specifically avoided contacting these colleagues, in order to base our conclusions on internationally published reports and not just personal communications. One alternative would have been to send questionnaires to treating clinicians around the world. Such as task is time-consuming, difficult and prone to selection biases; we know of no exhaustive list of infectious diseases experts in bone and joint infections. There is a Philadelphia Consensus (69) group that probably comes closest to what we imagine of such a community, but it is also composed of invited persons identified by their academic activity; without representing an exhaustive list of experts.

Conclusions

Based on the limited in vitro data for penetrating into bone, oral FLU could be seen as inferior to other oral agents that are not β-lactams, for which a bone penetration may be better. However, the role of antibiotic penetration, as commonly measured, in outcomes of treatment of bone infection is far from clear (70). Furthermore, most infectious diseases experts lack extensive personal experience in administering oral FLU as treatment for OM, while the small number who do (mostly from English-speaking or Northern European countries) have apparently not seen the need to publish their experience. The few published clinical studies clearly demonstrate that after any necessary debridement, a short course of intravenous FLU followed by oral FLU...
is a safe and effective option for treating OM (perhaps including DFO) in adults and children.

We encourage those who treat OM patients to consider conducting prospective trials, as treatment with oral FLU has several important advantages to make it worthwhile investigating in light of the problem of increasing antibiotic resistance: relatively low risk for adverse events and drug interactions; a narrow antibiotic spectrum; and, relatively low costs. Optimally, these studies would be randomized controlled trials comparing oral FLU to an oral quinolone. Statistically, the first option would probably require 2 x 232 OM episodes in a standard binary outcome non-inferiority trial with inferiority limits of 15%. Alternatively, even a prospective cohort study with oral FLU would be useful.

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Competing Interests

The authors have declared that no competing interest exists.

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