Management of Acute Native Joint Bacterial Arthritis in Adults in 2020 - A Short Narrative, Practical State-of-the-Art Review

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Originally published at:
Ansorge, A; Selman, F; Uçkay, İlker (2021). Management of Acute Native Joint Bacterial Arthritis in Adults in 2020 - A Short Narrative, Practical State-of-the-Art Review. Acta Scientific Orthopaedics, 4(1):54-59.
Management of Acute Native Joint Bacterial Arthritis in Adults in 2020 -
A Short Narrative, Practical State-of-the-Art Review

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Abstract

Native joint bacterial arthritis is a common infection among adults and children. A solely conservative management, without any articular drainage/lavage, increases the risk of recurrence. In contrast, the type of initial lavage/drainage can be surgical (arthrotomy or arthroscopy) or non-surgical (iterative arthrocenteses). Up to date, no superiority has been shown for any of these approaches in relation to recurrence risk and postinfectious mechanical damage. Furthermore, an initial synovectomy, or the number of iterative drainages does not influence outcome in most cases. Nowadays, an antibiotic regimen of three to four weeks, with early oral therapy, is standard in most settings of the world. In arthritis cases involving the hand and wrist, a shorter systemic antibiotic treatment such as two weeks is sufficient. The outcome of infection is impacted by of mechanical sequelae in up to 40% of cases. These sequelae are predominantly joint stiffness and/or osteoarthritis, which are difficult to treat.

Keywords: Native Arthritis; Antibiotics; Surgical Drainage; Management; Narrative Review

Introduction

The clinical appearance of native joint bacterial arthritis are very close to those of any non-infectious arthritis, such as viral, reactive, crystal-induced, or auto-immune arthritis. Yet, bacterial arthritis should be diagnosed and treated early in the process, since enzymes liberated by bacterial pathogens or reacting granulocytes risk to damage the bradytroph cartilage. Available literature about bacterial arthritis is composed of plenty expert opinions and reviews, while original research articles are less frequent [1]. In this narrative review, we focus on the current “lege artis” clinical management of native joint bacterial arthritis based on available published evidence.

Epidemiology, origins and causative pathogens

The annual incidence of native joint bacterial arthritis in wealthy countries varies between 2-10 cases per 100,000 persons [2-4] as well as patient charts with a discharge diagnosis of septic arthritis (International Statistical Classification of Diseases and Related Health Problems (ICD). Most septic arthritis is monoarticular: Indeed, only 2.7 to 15% present as a polyarticular arthritis [5-10] the proportion of septic arthritis cases involving people who inject drugs (PWID). The majority of septic arthritis cases occur spontaneously or are believed to be secondary to an hematogenous spread of bacteria from a skin lesion into the joint [2-4] as well as patient charts with a discharge diagnosis of septic arthritis (International Statistical Classification of Diseases and Related Health Problems (ICD). Spontaneous infection involves predominantly the major weight-bearing joints. Indeed, the knee is affected most of the time, followed in decreasing frequency by the hip, the elbow, the hand, the ankle, the wrist, the sternoclavicular and finally the sacroiliac articulations [6]. Globally, Staphylococcus aureus is the most preva...
lent pathogen in 50-60%, followed by streptococci (16%-17%), gram-negative rods (5%-15%), and anaerobes (<1%) [4,5] clinical pattern and outcome. We reviewed the literature from 1945 to 2010 with an emphasis on post-traumatic cases. We retrieved 14 large-scale epidemiological surveys without detailed stratification regarding the origin of septic arthritis (3,340 episodes). In contrast to spontaneous adult cases where Staphylococcus aureus is most prevalent, in infants the most encountered bacteria is a gram-negative rod named Kingella kingae [1,11] we aimed to contrast the bacteriologic epidemiology of osteoarticular infections (OAIs).

Diagnosis

Acute native joint arthritis usually presents with a swollen articulation, local erythema and heat, and worsening arthralgia. Most patients are transiently febrile. High fever spikes are uncommon, unless there is a synchronous bacteraemia [12]. One should in particular look for indices of remote infections like for instance endocarditis, respiratory, urinary tract, sexually-transmitted and skin infections. In sexually-active patients without adequate protection, Neisseria gonorrhoeae, is still quite prevalent [13]. Risk factors such as autoimmune rheumatic disease and intravenous drug abuse should also be looked for. Interestingly, patients infected with the human immunodeficiency virus don’t have an increased risk to develop septic arthritis unless they are concomitant intravenous drug abusers [1]. If the detailed anamnesis and the general physical exam fail to identify a remote origin of the septic arthritis, we generally don’t recommend to perform advanced diagnostic tests to force the recognition of the origin of the infection because the antibiotic treatment usually cure it, so the expenses for endoscopy and complex radiologic exams can be avoided [14]. One should also be aware that septic arthritis may coincide with crystalline arthritis [15,16]227 with underlying arthroplasties and 69 with gout or other crystals in synovial fluid. In such cases, it remains unclear if the crystalline inflammation is prejudicial in the acute phase. However, evidence exists in favor of the use of anti-crystalline treatment like colchicine in case of an acute gout developing just after the drainage of a joint for septic arthritis [1]. The gold standard for diagnosing bacterial arthritis is the objectivation of an identical pathogen in at least two samples of synovial fluid or intraarticular tissue biopsy, or a clinical arthritis associated with synchronous positive blood cultures that aren’t considered as contamination. Blood cultures should be performed in case of fever and shivering, considering that 50% of them turn positive [2].

However, a culture-negative bacterial arthritis may also occur: Risk factors for culture-negative cases are a low inoculum, a prior antibiotic exposure, an inappropriate culture media, fastidious bacteria, or a prolonged transport time to the laboratory favor negative-culture results [17]1167 (43%). A clear-cut definition of culture-negative bacterial arthritis doesn’t exist. It is usually diagnosed in cases showing a typical clinical presentation with suggestive pathologic synovial fluid laboratory results (granulocyte proportion of >90% and/or leukocyte count over 50'000 cells/µl), and no intraarticular crystals. Polymerase chain techniques (PCR) shouldn’t be routinely used, because eubacterial PCR is quite expensive and seldom of benefit. Its sensitivity is lower than that of classical culture. Moreover, it doesn’t give clues about antibiotic resistance and in the case of a polymicrobial infection its interpretation is hazardous [1]. In particular clinical situations suggesting an infection by one of the following pathogens, specific PCR is however recommended: Kingella kingae, Brucella spp, Borrelia spp, Coxiella burnetti, Neisseria gonorrhoeae, Bartonella henselae, Mycobacterium tuberculosis, or Mycobacterium ulcerans are such examples [1].

Laboratory inflammatory markers can be tested systemically (i.e. in the blood) and intraarticularly. High white blood cell counts and C-reactive protein concentrations in the blood serum aren’t proving a bacterial origin of an arthritis regardless of the used cut-off. Indeed, these inflammatory markers presumably rather reflect the pathogen’s virulence and the synchronous bacteraemia than the local intraarticular state [18,19]. Therefore, they are also useless in the follow up, concerning decision making about the duration of antibiotic treatment or the indication of repeated joint drainage/lavage. It is still important to know that serum procalcitonin concentration is often low in the case of a localized septic arthritis lacking systemic inflammatory signs [19,20]. Usually, synovial leukocyte counts and neutrophil proportions are determined. These tests must however be interpreted with caution, as no cut-off values have been shown to be diagnostic for bacterial arthritis. Concerning synovial leukocyte counts, many experts consider a number of >50,000 cells/µl as predictive for a bacterial arthritis [21]. However, there is equally no consensus about this threshold value and other authors set it for example at >100,000 cells/µl [22] not the surrounding soft tissues. The most common causes of monoarthritis are crystals (i.e., gout and pseudogout. Concerning the neutrophil proportion in the synovial fluid, a literature review including 6242 cases showed a three-fold probability for septic arthritis in patients with a neutrophil proportion of >90% in comparison to
prompt identification and treatment of septic arthritis can substantially reduce morbidity and mortality. OBJECTIVE: To review the accuracy and precision of the clinical evaluation for the diagnosis of nongonococcal bacterial arthritis. DATA SOURCES: Structured PubMed and EMBASE searches (1966 through January 2007). In future, other intra-synovial marker tests may appear. For instance, research about intraarticular lactic acid dosing is warranted as well as in 12 cases of gonococcal arthritis (mean 27 mg/100 ml).

Classical direct microscopic exams using Gram and acridinorange staining have been abandoned by many centers as its cost-benefit analyses are unsatisfying. The low sensitivity of direct microscopy isn’t surprising as a low intraarticular inoculum is enough to trigger a significant arthritis. The results of Cunningham, et al. illustrate well the low sensitivity of direct microscopy with underlying arthroplasties and 69 with gout or other crystals in synovial fluid. In their series of 500 adult patients with suspected septic arthritis, they detected pathogens in only 29% of cases. When considering only the culture-positive cases (gold standard), they still yielded a low sensitivity of 37%.

Therapy

The treatment of native joint septic arthritis in adults is mainly based on experts’ opinion and on regional habits, because evidence is lacking. As a consequence, the treatments vary among many parts of the world. However, it is generally articulating around two principles, namely lavage/drainage of the affected joint(s) and synchronous systemic antibiotic therapy. Available evidence suggests a higher recurrence risk if only one of these two principles is used for treatment. For instance, a retrospective work found a 21-fold increase of the recurrence risk in case of a strict conservative antibiotic management without any lavage/drainage. A strict surgical management with iterative lavages/drainages is in turn impacted by a higher probability of deformity and consecutive osteitis.

Drainage

Acute bacterial arthritis is usually considered as an absolute emergency, necessitating immediate lavage/drainage of the affected joint. This opinion is relying basically on animal studies, while available clinical studies in humans fail to support it caused by autoimmunity, immune complexes, crystals, or cartilage damage, is clinically similar to native septic arthritis. The large microbiological spectrum of joint infection is one of the particularities distinguishing arthritis from other orthopedic infections such as osteomyelitis or implant-related infections. The current literature often defines orthopedic infections as "osteoarticular," suggesting that diagnosis and treatment of bone and joint infection would be similar. Septic arthritis, osteomyelitis, and orthopedic implant–associated infections are different in nature, epidemiology, therapy, and outcome. This chapter underlines the particularities of native joint arthritis with an emphasis on diagnosis, epidemiology, treatment, and some prevention aspects before planned joint interventions. Most cases of primary septic arthritis and surgical site infections (SSIs. For instance, Lauper, et al. retrospectively analyzed the long-term outcome of over 200 native joint adult bacterial arthritis cases in relation to the delay of surgical drainage since the hospital admission requiring drainage within hours, including during night, weekend or holiday shifts. However, there are few data supporting the need for the disruption caused by this degree of urgency. METHODS: We performed a retrospective review of all adult patients seen in our medical center from 1997-2015 with culture-proven septic arthritis and noted the epidemiology of sequelae, and their possible association with a delay in surgical drainage. RESULTS: Of 204 septic arthritis episodes, 46 (23%). Interestingly, no outcome difference could be shown between a delay of less than 6 hours, 6-12 hours or even exceeding 24 hours. They also found a mean span of symptoms of 3 days ahead of hospital admission. Another study by Vispo-Seara, et al. found a timeline between the onset of septic arthritis and arthroscopic lavage/drainage of over 2 weeks as being associated with advanced cartilage damage. Balabaud, et al. have found a delay of less than 12 days until surgical drainage to be associated with healing of knee septic arthritis, as compared with a delay of 23 days associated with treatment failure.

Current literature indicate that any type of joint drainage is valid. The choice of drainage type is also depending on regional practice. While surgeons prefer arthrotomy or arthroscopy for drainage, many rheumatologists choose iterative arthrocentesis.
Concerning adult septic arthritis, no randomized clinical trials comparing surgical drainage to iterative arthrocentesis are available. Only retrospective reports are existing. For instance, Harada, et al. conducted a ten year retrospective single-center study to compare outcomes based on the taken management approach: medical (bedside closed-needle joint aspiration) versus surgical (arthrotomy/arthroscopy) [29] based on medical versus surgical management.

METHODS: A 10-year retrospective single-center study was conducted of patients admitted to a tertiary care hospital between January 1, 2006 and December 31, 2015 with a diagnosis of SA to compare outcomes based on the management approach taken: medical (bedside closed-needle joint aspiration). They evaluated outcomes were: the joint recovery, time to recovery, length of hospital stay, disposition to home versus rehabilitation unit, recurrence of arthritis in the same joint, and mortality.

There was no statistically significant difference in long-term outcomes between the two groups at 12 months. Some surgeons plan a second look after the initial surgical drainage/lavage, based exclusively on the intraoperative visual aspect of the joint or on the fact that serum C-reactive protein concentrations aren’t decreasing in further follow-up. This attitude isn’t supported by evidence. We rather recommend to base the indication for a repeated surgical drainage according to the global clinical course.

Antibiotic treatment

The spectrum

Until the susceptibility results are available, the initial empirical antibiotic therapy in bacterial joint arthritis should cover the most frequent pathogens. After that, the initial regimen should be replaced by the narrowest possible spectrum. Methicillin-susceptible Staphylococcus aureus and streptococci should be covered at the beginning. In case of posttraumatic arthritis (e.g., bites or splinter injuries), Gram-negative bacteria have to be covered as well. In such cases, empirical intravenous (IV) administration of first- or second-generation cephalosporin or aminopenicillin is indicated. If community-acquired MRSA (Methicillin-resistant S. aureus) is suspected, vancomycin is indicated; respectively clindamycin or daptomycin, if an allergy to β-lactam antibiotics is present [32]. According to the expert opinion, empirical therapy starts intravenously [1]. All intravenous and the majority of oral antibiotics do penetrate enough the synovia and do reach concentrations exceeding the minimal inhibitory breakpoints of common pathogens [33,34]. In acute virulent arthritis early IV-application makes sense, but a switch to moral therapy is possible as soon as the clinical course is favorable. All non-beta-lactam antibiotics, as for example cotrimoxazole and quinolones are well-known for their good oral synovial bioavailability [35,36] where, for at least the first 2-6 weeks, antibiotics should be administered intravenously, is more and more challenged in favor of an oral antibiotic treatment with selected agents from the start. There is no evidence that the total duration of antibiotic therapy for more than 4-6 weeks improves outcome, when compared with shorter regimens.

Hopefully, the future will show randomized trials in the adult population, allowing optimal timing of surgical and medical therapy and sparing of unnecessary prescription, with concomitant development of antibiotic resistance. External advice from an expert team with combined surgeons and infectious disease physicians may help to reduce antibiotic consumption in a cost-effective way. [1,29-31].

Citation: Ilker Uçkay. "Management of Acute Native Joint Bacterial Arthritis in Adults in 2020 - A Short Narrative, Practical State-of-the-Art Review". Acta Scientific Orthopaedics 4.1 (2021):.
treated with antiseptics partly followed by irrigation with sodium chloride solution was performed by using Casy Cell-Counter. Light microscopic data revealed a defect in cell structure after addition of antiseptics. We showed a significant increase of LDH enzyme activity after the treatment with polyhexanide or taurolidine. After treatment with antiseptics followed by sodium chloride solution a significant increase of vital and total cell numbers resulted in comparison with the chondrocytes that were only treated with antiseptics. The data show that treatment with polyhexanide, hydrogen peroxide or taurolidine induces cell death of human chondrocytes in vitro. The application of sodium chloride solution after the treatment with polyhexanide and hydrogen peroxide possibly has a protective effect on chondrocyte viability.

Toxicity of antiseptics against chondrocytes: What is best for the cartilage in septic joint surgery? The optimal duration of post-drainage systemic antibiotic treatment is controversial. Often the indicated therapy might depend on the individual case. Some experts advocate a 3-4 weeks IV antibiotic therapy for staphylococci and Gram-negative pathogens, a 2-week IV therapy for streptococci, and an IV therapy exceeding four weeks for immune-compromised patients [2]. Nade., et al. recommend a 2 weeks IV therapy for streptococci [38]. Often, an IV therapy is recommended for the first 2 weeks, followed by a 2-week oral therapy. Yet, this duration is currently considered as excessive for drained, adult native joint infections. Indeed, a total antibiotic treatment duration of 3-4 weeks is sufficient [33,39]. The antimicrobial treatment in bacterial hand and wrist arthritis is usually very short. Angly., et al. showed that no recurrence occurred in operated adult finger arthritis after antibiotics administered for a median duration of 2 days IV and 17 days orally [40].

Duration of antimicrobial therapy

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Outcomes

Primary bacterial native joint arthritis is exceptionally fatal. If it is so, usually a severe underlying remote origin, such as endocard-
dritis or a concomitant sepsis is usually responsible for the mortal outcome. Indeed, local infection remission frequently occurs in 97% in small joints, respectively in 90-95% in large joints [18,42] to gather data for a prospective study on an optimized antibiotic treatment in adults with septic arthritis.

METHODS: This was a retrospective single-center study conducted for the period 1996-2008. RESULTS: A total of 169 episodes of septic arthritis in 157 adult patients (median age 63 years; 65 females. In contrast to the good infection control after eventually several drainages, the mechanical consequences are of major concern. Prospective randomized trials in adults with septic arthritis reported a frequency of mechanical sequelae between 20% and 35%, of which 15% required further surgical intervention [26]. METHODS: We performed a retrospective review of all adult patients seen in our medical center from 1997-2015 with culture-proven septic arthritis and noted the epidemiology of sequelae, and their possible association with a delay in surgical drainage. RESULTS: Of 204 septic arthritis episodes, 46 (23%). Currently efforts are done to try to treat them by re-education, sensory-integrative therapy and eventual corrective surgery such as joint prostheses.

Conclusion
Native joint bacterial arthritis is common. An articular drainage/lavage (arthrotomy or arthroscopy, or iterative arthrocentesis) is always warranted. Up to date, no superiority has been shown for any of these approaches in relation to recurrence risk and postinfectious mechanical damage. Furthermore, an antibiotic regimen of three to four weeks, with early oral therapy, is standard. In arthritis cases involving the hand and wrist, a shorter systemic antibiotic treatment such as two weeks is sufficient. The outcome of infection is impacted by of mechanical sequelae, which are difficult to treat and need farther research for clinical improvement.

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