Osteoarticular infection in intravenous drug abusers: influence of HIV infection and differences with non drug abusers


Abstract

Objectives—To determine (a) the influence of HIV in developing osteoarticular infections in intravenous drug abusers (IVDAs) and (b) the differences between the clinical features of osteoarticular infections in IVDAs and a control group of non-IVDAs.

Methods—A comparative study of the clinical features of osteoarticular infections in all HIV positive and HIV negative IVDAs admitted to the departments of rheumatology and internal medicine during a 10 year period was carried out. The joint infections of all IVDAs, irrespective of HIV status, were compared with those of a control group of non-IVDAs lacking risk factors for HIV infection.

Results—A total of 482 HIV positive and 85 HIV negative IVDAs was studied, in whom 25 (5%) and six (7%) osteoarticular infections were found respectively. There were no differences in age, sex, joints affected, and causative agents between these two groups. A comparison of the 31 (5.5%) osteoarticular infections in all IVDAs with 21 infections in 616 (3.4%) non-IVDAs showed significant differences in the mean age (27.5 v 54), the frequency of affection of the axial joints (hip, sacroiliac, and sternocostal joints) (64.5% v 16–6%), and in the incidence of Candida albicans (19% v 0%).

Conclusions—(1) HIV may not predispose to osteoarticular infections in IVDAs. (2) The hip, sacroiliac, and sternocostal joints (axial joints) were most commonly affected in IVDAs. (3) In Spain, unlike other countries, Gram positive bacteria and C albicans seem to be predominant agents in osteoarticular infections in IVDAs, with a low incidence of Gram negative bacteria.

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The human immunodeficiency virus (HIV), one of the main protagonists of the last decade, has a role in all clinical fields. In 1985 psoriasis was first reported in relation to HIV,1 and in 1987 Reiter's syndrome was described as a possible HIV associated disease.2 Since these initial reports some investigators have studied the incidence and prevalence of rheumatic syndromes in association with HIV infection. A predominance of seronegative spondyloarthritides in studies among homosexual men3–5 and of osteoarticular infections among intravenous drug abusers (IVDAs)6–10 may exist. Furthermore, septic arthritis has been found only rarely when homosexuality is the principal risk factor for HIV infection.3–5 This suggests that risk factors for HIV infection may influence the expression of HIV related rheumatic syndromes.

On the other hand, septic arthritis in IVDAs might have different features than in non-IVDAs. The former might have infections of the axial joints more commonly than the latter; this has not been proved.

The objectives of our study were to determine (a) the influence of HIV on the incidence of osteoarticular infections in IVDAs and (b) the differences between osteoarticular infections in IVDAs and non-IVDAs.

Patients and methods

We included in the study all IVDAs (with or without HIV infection) admitted to the department of internal medicine and rheumatology of our hospital from 1981 to 1990. Patients with other risks for HIV infection were excluded. In all patients a clinical protocol was completed, which included age, sex, joints affected, causative agent and site of isolation, predisposing factor for osteoarticular infection, HIV status and group of Centers of Disease Control (CDC) classification in positive cases, radiological abnormalities, treatment given, and response achieved. A possible relation between acquired immunodeficiency syndrome (AIDS) and osteoarticular infections was specifically looked for. Diagnosis of an episode of osteoarticular infection was made by a compatible clinical picture of septic arthritis, discitis, or osteomyelitis and isolation of the causative agent in synovial fluid, blood, or tissue biopsy specimen.

Intravenous drug abusers were divided into HIV positive and negative groups. HIV serological tests were performed by an enzyme linked immunosorbent assay (ELISA) and confirmed by western blot. AIDS was diagnosed according to the CDC criteria as previously published.11 HIV infected patients were classified by the CDC system.12 The presence of osteoarticular infections, age, causative agents, and joints affected were compared in the two groups.
To investigate a possible relation between osteoarticular infections and a more profound immunological depression we subdivided HIV positive patients with osteoarticular infections in accordance with the CDC classification system (table 1). This distribution was compared with the prevalence of each CDC group in the HIV positive patients without osteoarticular infections.

To determine the differences between osteoarticular infections in IVDAs and non-IVDAs we compared age, predisposing factors for a skeletal infection, causative agents, and affected joints in all IVDAs, irrespective of their HIV status, with a group of non-IVDAs with osteoarticular infections. The non-IVDAs were recruited from all subjects lacking risk factors for HIV infection admitted to the department of rheumatology over five years (1986 to 1991).

Statistical analysis was performed with Fisher's exact test. A difference between the mean age of each group more than 2 or 2.6 times greater than the standard error was considered significant (p<0.05 or p<0.01 respectively). When the groups were small we compared the mean age using the appropriate correction of the standard error and Student's _t_ values.

**Results**

**Comparative study of IDVAs with and without HIV infection**

During a 10 year period 567 IVDAs were studied. Of these, 482 (85%) were HIV seropositive and 85 (15%) HIV seronegative. Mean age was 27 years in the HIV positive patients (range 17–39 years) and 29 in the HIV negative patients (range 23–37 years). In the HIV positive group 80% were men and 20% women, and the HIV negative group comprised 83% men and 17% women.

We found 25 osteoarticular infections (5%) in HIV positive patients and six (7%) in HIV negative patients. In the first group the affected joints were the hip (seven patients (28%)), sacroiliac joint (four (16%)), sternocostal joint (four (16%), knee (three (12%)), and shoulder and disc L3–4 in one case each. In one patient a polyarticular involvement of elbows and wrists was found, and in another the ankle and shoulder were both affected. Osteomyelitis was present in three patients (wrist, femoral neck, and tibia). In HIV negative patients the affected joints were the sacroiliac joint in three (50%) patients and hip, sternocostal joint, and knee in one each. Although sacroiliac joint infection was more common in HIV negative than in HIV positive patients (50% v 16%), the difference was not significant. Although osteomyelitis was not found in the HIV negative group, statistical analysis was not possible owing to the small number of HIV positive patients with osteomyelitis.

In the 25 HIV positive patients with osteoarticular infections the causative agents were _Staphylococcus aureus_ in 12 (48%), _Candida albicans_ in five (20%), and _Mucor, Neisseria gonorrhoeae, Mycobacterium tuberculosis, Staphylococcus epidermidis, Streptococcus viridans, Streptococcus equisimilis_, and _Streptococcus sanguis_ in one case each. Additionally, _S aureus_ and _Bacteroides melaninogenicus_ were found in one patient.

In the six HIV negative patients with osteoarticular infections the causative agents were _S aureus_ in four (67%) and _C albicans_ and _M tuberculosis_ in one case each.

No significant differences in the incidence of infections, age, sex, joints affected, and causative agents were found between HIV positive and negative patients (table 2).

Of the 25 HIV positive patients with osteoarticular infections, 11 (44%) were in CDC group II, four (16%) in group III, four (16%) in group IV–C2, and six (24%) in group IV–Cl and had criteria of AIDS.

### Comparative study of IVDAs and non-IVDAs

A group of 616 non-IVDAs with no risk factor for HIV infection were compared with all the IVDAs, irrespective of HIV status. In the former group 21 (3.4%) had osteoarticular infections and in the latter 31 (5.5%). The mean age of the non-IVDAs was 54 years (range 15–78) and of the IVDAs 27 ± 5 years (range 17–39), which was significantly different (p<0.001). Of the 21 non-IVDAs, 13 (62%) were men and eight (38%) women, whereas of the 31 IVDAs, 25 (81%) were men and six (19%) women.

In 16 of the 21 non-IVDAs with osteoarticular infections a predisposing factor was found. Six had previous skin or wound infections, and rheumatoid arthritis, diabetes
mellitus, pneumonia, and alcoholism were each present in two patients, one had systemic lupus erythematosus, and another had multiple myeloma. In addition to the soft tissue and skin infections commonly found in IVDAs, six of the 31 with osteoarticular infections (20%) had disseminated candidiasis (with the sternocostal joint affected in five patients and the hip in one) and six (20%) had infectious endocarditis. Of these, five had tricuspid endocarditis caused by S aureus and one had mitral endocarditis due to S sanguis. (This patient also had osteomyelitis of the tibia.)

The affected joints in the 21 non-IVDAs were knees (eight patients (38%)), intervertebral discs (three (14%)), wrists (two (10%)), elbows (two (10%), hips (two (10%)), and ankle, sacroiliac joint, and shoulder in one case each. Another patient had septic oligoarthritis of the wrist, shoulder, and ankle. Classifying the affected joints as axial (sacroiliac, sacroiliac, intervertebral, and hip joints) or peripheral, we found that the axial joints were more commonly affected in the IVDAs than in the non-IVDAs (64.5% vs 16.6%; p=0.00034). The knee was most commonly affected in non-IVDAs (eight v four cases), but this difference was not significant when compared with the IVDAs.

The bacteria responsible for the osteoarticular infections in the non-IVDAs were S aureus (67%), M tuberculosis (four (19%)), S pneumoniae (two (10%)), and B fragilis in one case (5%). The only significant difference found between IVDAs and non-IVDAs was the prevalence of C albicans (five v 0; p=0.03).

**Discussion**

**INFLUENCE OF HIV IN DEVELOPING OSTEARTICULAR INFECTIONS**

Since 1988 at least six large series have been published describing rheumatic diseases in HIV infected patients.3 5 7 9 10 In those studies with a high proportion of homosexual men (75–92% of HIV infected patients) a large number had seronegative spondylarthritides, mainly Reiter's syndrome, and no osteoarticular infections.3 5 In those studies with a high number of IVDAs (more than 80%), however, fewer had spondylarthritides and osteoarticular infections predominated.7 9 10 Hence the group at risk for HIV infection might influence the expression of HIV related rheumatic syndromes.

If the latter studies are excluded, osteoarticular infections in association with HIV infection are rarely reported. The first report published in 1985 described arthritis of small joints of the hand and wrist caused by *Sporothrix schenckii.*13 Since this report other infections in HIV positive patients caused by cryptococcus,14 S aureus,15-17 and C albicans18 have been reported. Many of these occurred in association with disseminated infections. We published a preliminary description of articular mucormycosis included in this study.19 At least six cases of septic arthritis (one salmonella, two pneumococcus, and three *S aureus*) occurred in six HIV positive haemophiliac patients.20 21 Nonetheless, infectious arthritis can be a complication in HIV negative haemophiliac patients.

A recent publication22 reported 10 episodes of musculoskeletal infections in nine patients from a group of about 3000 HIV positive patients (0.3%). Of the nine patients, eight were homosexual and one was an IVDA. Five of them had septic arthritis, two had septic arthritis with osteomyelitis, one had osteomyelitis, one had muscular abscess, and one had bursitis. The infections were caused by *S aureus* in four, *Salmonella enteritidis* in two, and pseudomonas, *Histoplasma capsulatum,* gonococcus, and *M kansas* in one case each. Of the nine cases, seven had criteria for AIDS.

To our knowledge this is the first study in which the role of HIV in the development of osteoarticular infections has been examined. Our data showed no differences between HIV positive and HIV negative IVDAs for mean age, sex, affected joints, and causative agents. This suggests that HIV may not play a part in the development of osteoarticular infections. Possibly, the results were biased as the group of HIV negative patients was small. Thus further studies are needed.

Most of our HIV positive patients with osteoarticular infections were in CDC groups II and III, and only six of the 25 (24%) were in group IV–Cl. Therefore profound immunological depression was present in only a few cases.

**DIFFERENCES BETWEEN OSTEARTICULAR INFECTIONS IN IVDAS AND NON-IVDAS**

About a half of IVDAs will develop an infectious complication,23 of which osteoarticular infections comprise 3–4%–4%.23 24 From 1983 to 1988 the 'Spanish working group for the study of infections in drug addicts' registered 11645 infections, and septic arthritis or osteomyelitis was present in 291 (2.5%).25 Sacroiliac joints were most commonly affected (23%), and *S aureus* was the most commonly isolated agent (55%).

Sacroiliac, sternocostal and sternocostal joints are common sites of septic arthritis in IVDAs (table 3).26-28 These sites, however, are atypical in non-IVDAs,30 in whom the knee, shoulder, wrist, hip, interphalangeal, and elbow joints are the typical localities.31-34 As for IVDAs, *S aureus* is the most common cause of infection in non-IVDAs,31-34 and candida is almost never present.

Although there are some differences between osteoarticular infections in IVDAs and non-IVDAs, we found no studies in which these differences were specifically examined. Our paper confirms that in IVDAs a significantly greater number of infections of axial joints—namely, hip, sacroiliac, and sternocostal joints—are present than in non-IVDAs. In addition, *C albicans* is found significantly more often in IVDAs than in non-IVDAs.

All the non-IVDAs studied were hospital inpatients and, furthermore, almost all (76%) had a predisposing factor for joint infection. Thus the incidence of osteoarticular infections
(3.4%) in this group cannot be extrapolated to the general population.

Intravenous drug abuse, itself, can be considered a risk factor for articular infections. Ang-Fonte et al attribute a fourfold increase of septic arthritis to this factor.35 In his paper 75% of 28 patients with non-gonococcal arthritis evaluated during 1981 and 1982 were IVDAs. This is a very high percentage when compared with the incidence of 35% reported in the same hospital during 1966–77.

The causative agents responsible for osteoarticular infections are similar to those found by others in Spain.36 29 In comparison with available American data, we and other Spanish authors found a lower number of Gram negative bacteria and methicillin resistant S aureus. This suggests that geographical variations in the epidemiology of such infections may occur. It should be emphasised that the incidence of articular tuberculosis was not increased in IVDAs.

Several factors may account for the high incidence of infections in IVDAs: (a) HIV infection might be an important factor; but our data suggest that in osteoarticular infections this virus plays little part; (b) S aureus, the most common cause of infection, has not been found in heroin samples, but IVDAs are often carriers of S aureus; (c) unsterile drugs are injected with dirty needles and syringes; (d) dental hygiene is often poor; (e) bacterial clearance by the tracheobronchial system can be decreased during intoxication; (f) promiscuity and prostitution increase the risk of transmitting infective agents; and, finally, (g) cell mediated immunity may be impaired in IVDAs not only by HIV infection but also by intravenous drugs themselves.37

It is known that infectious agents each reach the synovial space through the bloodstream.31 Why do IVDAs have a greater number of infections in the axial joints? Perhaps, the site of injection may play a part.27 This hypothesis, however, does not completely explain the increase of infections of sacroiliac, hip, and other axial joints in IVDAs. The high incidence of sternocostal infections is explained by disseminated candidiasis, which was present in 20% of our IVDAs. We found five cases of sternocostal infection and all were caused by this agent. The high incidence of C albicans in IVDAs seems to be due to the use of brown heroin mixed with contaminated lemon.38 39

Therefore, the presence of disseminated candidiasis does not imply immune suppression related to AIDS. Systemic candidiasis has an initial phase with a septicemica febrile syndrome that lasts between one and three days and a second phase characterised by the appearance of metastatic lesions, such as cutaneous (folliculitis, nodules, etc), ocular, and articular disease. The time between the acute phase and the articular disease ranges from 15 days to five months,38 and sternocostal infection is the most common and characteristic joint disease.39 The reason for this is unknown.

In conclusion, after searching for the influence of HIV infection in developing osteoarticular infections in IVDAs, we found that HIV seems to play little part. Intravenous drug abusers are younger and have a greater number of axial joint infections and have more arthritides due to candida than non-IVDAs. The epidemiology of the infections in our patients, similar to that reported by others in Spain, shows less Gram negative bacterial infections and methicillin resistant S aureus than in the United States.

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Table 3 Clinical features of the main reports about septic arthritis in drug addicts

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<tr>
<th>Reference</th>
<th>Affected joints</th>
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