A Retrospective Analysis of Drug Fever Diagnosed during Infectious Disease Consultation

Kenichiro Yaita, Yoshiro Sakai, Kenji Masunaga and Hiroshi Watanabe

Abstract

Objective To clarify the current situation concerning drug fever (DF) in Japan, we retrospectively analyzed patients undergoing infectious disease consultation at our institution.

Methods Between April 2014 and May 2015, we extracted the records of DF patients from among 388 patients who had obtained infectious disease consultations in Kurume University Hospital. We reviewed their medical charts and summarized the characteristics of DF.

Results This study included the records of 16 patients. Clinical signs (relative bradycardia, the duration of the drug administration before becoming febrile, and the interval between the discontinuation of a drug and the alleviation of a fever), and laboratory tests (varied white blood cell count, low level of C-reactive protein, and a mild elevation of transaminases) were compatible with those from previous reports. Among the drug-confirmed cases, five involved the use of glycopeptides (vancomycin: 3, teicoplanin: 2), which were considered to be uncommon causes, and the another five cases involved the use of β-lactams. In addition, the procalcitonin levels were either negative or low (<0.25 ng/mL) in 10 of the 11 procalcitonin-measured cases.

Conclusion Our findings demonstrated that glycopeptides, similar to β-lactams, may be the origin of DF. Furthermore, procalcitonin may be helpful in the diagnosis of DF, but only in combination with other detailed examinations.

Key words: drug fever, glycopeptides, vancomycin, teicoplanin, procalcitonin


Introduction

We occasionally encounter patients who are febrile, yet non-infectious, due to drugs. Although the mechanism of drug fever (DF) is unclear, the most common reason is thought to be the result of immune-mediated/hypersensitivity reactions (1). The typical characteristics of DF include an onset 7 to 10 days after drug administration, relative well-being, relative bradycardia, varied white blood cell (WBC) count (including neutropenia), elevation of eosinophils, and mildly elevated serum transaminases (1-4). Although these signs are well known, the diagnosis of DF remains challenging, because the correct diagnosis requires the exclusion of other non-localized febrile illnesses such as bacteremia. Eventually clinicians diagnose DF by a prompt improvement (within 3 days) after discontinuation of the suspected drug(s) (1).

Antibiotics are thought to be the most common offending drugs (3, 5), and β-lactams and trimethoprim/sulfamethoxazole are predominantly fever-causing antibiotics (2). A series of case reports concerning DF have been reported in the U.S., France, and Japan (6-11). In most of these reports, β-lactams are the common offending drugs, and the clinical characteristics of these DF cases were also compatible with previous review articles.

However, there has been very little new information concerning DF in Japan. The latest reports of a Japanese DF case series (10, 11) were published more than 20 years ago. We herein present an analysis of DF cases through the experience of infectious disease (ID) consultations in order to clarify the current tendencies of DF in Japan.
Table 1. The Characteristics of Patients with Drug Fever.

<table>
<thead>
<tr>
<th>(n=16)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean ± SD (year)</td>
</tr>
<tr>
<td>Sex (male)</td>
</tr>
<tr>
<td>Past drug allergies a</td>
</tr>
<tr>
<td>Primary illnesses</td>
</tr>
<tr>
<td>Bacteremia (including IE)</td>
</tr>
<tr>
<td>Postoperative fever after TAR (allergic reaction to vascular graft, suspected)</td>
</tr>
<tr>
<td>Pneumonia (HCAP and VAP)</td>
</tr>
<tr>
<td>Infective aortic aneurysm (suspected)</td>
</tr>
<tr>
<td>Surgical site infection (suspected)</td>
</tr>
<tr>
<td>Postoperative intra-abdominal abscess</td>
</tr>
<tr>
<td>Febrile neutropenia</td>
</tr>
</tbody>
</table>

a: Includes a suspected case. b: Includes a patient that underwent an operation for an open fracture of the lower leg and a patient that underwent an operation for funnel chest. HCAP: health-care associated pneumonia, IE: infective endocarditis, SD: standard deviation, TAR: total arch replacement, VAP: ventilator-associated pneumonia

Table 2. The Clinical Characteristics of Patients with Drug Fever.

<table>
<thead>
<tr>
<th>(n=16)</th>
</tr>
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<tbody>
<tr>
<td>Fever-causing drugs</td>
</tr>
<tr>
<td>Vancomycin</td>
</tr>
<tr>
<td>Teicoplanin</td>
</tr>
<tr>
<td>Piperacillin/Tazobactam</td>
</tr>
<tr>
<td>Ampicillin/Sulbactam</td>
</tr>
<tr>
<td>Cefazolin</td>
</tr>
<tr>
<td>Ceftriaxone</td>
</tr>
<tr>
<td>Not clarified because two suspected drugs were discontinued at the same time</td>
</tr>
<tr>
<td>The duration of the drug administration before becoming febrile, mean ± SD (days)</td>
</tr>
<tr>
<td>The interval between the discontinuation of the drug and alleviation of the fever, mean ± SD (days) b</td>
</tr>
<tr>
<td>The type of fever</td>
</tr>
<tr>
<td>A gradual increase</td>
</tr>
<tr>
<td>Spike</td>
</tr>
<tr>
<td>Eruption</td>
</tr>
<tr>
<td>Relative bradycardia b</td>
</tr>
<tr>
<td>Highest BT, mean ± SD (ºC)</td>
</tr>
</tbody>
</table>

a: Two patients were excluded because the duration of the administration of two suspected drugs were different. b: Includes three patients taking β-blockers and one patient taking calcium-channel blockers (amlodipine and azelnidipine). BT: body temperature, SD: standard deviation

Materials and Methods

This research involved a single-center study that was conducted by the Department of Infection Control and Prevention, which manages ID consultations for every clinical department in Kurume University Hospital, a 1,025-bed tertiary care medical center. The Kurume University Research Ethics Committee (http://www.med.kurume-u.ac.jp/med/joint/irnr/ir) approved this study (Research No. 14267). Between April 2014 and May 2015, 388 patients underwent ID consultation. We retrospectively reviewed their medical charts, and extracted the information of DF patients.

DF was defined by clinical characteristics that met all of the following criteria: 1, an axial temperature above 37.5°C (in Japan, an oral or rectal temperature is generally not measured); 2, no other origin of fever can be detected by detailed ID consultation (including appropriate imaging tests and microbiological tests); 3, any underlying febrile illness, the improvement of which can be confirmed by the ID physician; and, 4, after the discontinuation of drugs, the fever is alleviated. These criteria were made in reference to previous studies (6-11) with some modification. We classified the following factors as patient-related: age, sex, primary illnesses, known allergies, antibiotics use, the duration of the drug administration before becoming febrile (days), clinical symptoms [the type of fever, the highest body temperature, eruption, and relative bradycardia (as defined by Cunha’s report (2))], laboratory findings [peak WBC count, including neutropenia, and the levels of C-reactive protein (CRP), aspartate aminotransferase (AST), alanine aminotransferase (ALT) and procalcitonin (PCT)] during the febrile period, and the interval between the discontinuation of a drug and the alleviation of the fever (days).

Results

Among the 388 ID consulted cases, 16 patients met all the qualifications for inclusion into this study. We could not determine the fever-causing agent in 6 cases because two drugs had been simultaneously discontinued. The profiles of the DF cases are summarized in Table 1. The mean age ± standard deviation (SD) was 63.6±19.4 years, and 10 patients were men. The primary illness of 5 of the patients was bacteremia with or without infective endocarditis. Two patients had allergies including one suspected case. One patient had a previous history of drug eruption due to cefoperazone/sulbactam, and another patient had a drug fever suspected history during the administration of meropenem and vancomycin. The clinical characteristics of patients with DF are described in Table 2. In the drug-confirmed cases, 5 patients had used glycopeptides (vancomycin: 3, and teicoplanin: 2), and the other 5 cases were caused by β-lactams (piperacillin/tazobactam: 2, ampicillin/sulbactam: 1, cefazolin: 1, and ceftriaxone: 1). The antibiotic therapies were discontinued in 6 cases, while the fever-causing drugs in 10 cases were discontinued and changed to other drugs. The mean duration of the drug administration before becoming febrile averaged 8.6±5.3 days, and the mean interval between the discontinuation of the drug and the alleviation of the fever averaged 3.4±3.3 days. Clinical signs such as a gradually increasing fever, eruption and relative bradycardia were seen in 10, 5, and 14 patients, respectively, and the mean highest body temperature was 38.8±0.8°C. The laboratory test results of patients with DF are presented in Table 3. The laboratory tests showed a peak WBC count of 7,519±3,551/µL (eosinophil count 429±575/µL), a peak CRP level of 5.1±3.9 mg/dL, a peak AST level of 47.7±46.0 IU/L, a peak ALT level of 48.9±59.5 IU/L, and PCT was negative in 8 patients. During febrile periods, neutropenia and eosinophilia were detected in 4 and 4 patients, respectively.
resistant and frequently used drugs. The ratio of methicillin-
multi-drug resistant bacteria, glycopeptides have become im-
endocarditis (15). In the modern era, with its prevalence of
infections (14); and another study described 5/20 cases that
the treatment of bone, joint, and vascular-access-associated
DF and rash (13). Other reports have included DF due to
their study reported 24 cases of DF alone and 8 cases of
artery (9, 13-15). In a French national cohort study, which is
previous cases have been described in the litera-
ture (5), which could have resulted in a bias in the
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triggers for suspecting DF.

None of the patients exhibited serious drug-reactions.

### Discussion

In this report, characteristics such as the duration of drug
administration before becoming febrile, the time until im-
provement after the discontinuation of the drugs, relative
bradycardia, a moderate to slightly elevated WBC count, a
low level of CRP, and a mild elevation of transaminases
were compatible with the results shown in previous reports.
However, the prevalence of relative bradycardia and glyco-
peptides use was much higher than we expected. Further-
more, the PCT levels in most patients were either negative
or low. All of our participants were seen for ID consultation,
and relative bradycardia is an important clinical clue for ID
physicians (5), which could have resulted in a bias in the
triggers for suspecting DF.

Although glycopeptides, including vancomycin and teico-
planin, are considered to be rare DF-causing agents (12),
previous cases have been described in the litera-
ture (9, 13-15). In a French national cohort study, which is
the most recent report concerning DF, there were 5 DF
cases caused by glycopeptides from among 167 DF pa-
patients (9). Hung et al. described 117 cases where the medi-
cation was switched from vancomycin to teicoplanin due to
DF, rash or neutropenia during the treatment period (13). Their
study reported 24 cases of DF alone and 8 cases of
DF and rash (13). Other reports have included DF due to
teicoplanin; one study reported 5/18 cases with rash during
the treatment of bone, joint, and vascular-access-associated
infections (14); and another study described 5/20 cases that
included 3 with rash during the treatment of streptococcal
endocarditis (15). In the modern era, with its prevalence of
multi-drug resistant bacteria, glycopeptides have become im-
portant and frequently used drugs. The ratio of methicillin-
resistant Staphylococcus aureus / methicillin-susceptible
Staphylococcus aureus sampled from clinical specimens in
our hospital in 2014 was 1.07 (1,334/1,243 samples) (un-
published data), which may have been associated with the
frequent use of glycopeptides, and there was a high preva-
ence of glycopeptides as the causative agents of DF in the
present study. Although the characteristics of glycopepti-
duced DF patients were similar to those of the patients af-
fected by other drugs, one patient had a persistent fever for
12 days after the discontinuation of the teicoplanin. Piper-
acillin/tazobactam was administered at the surgeon’s direc-
tion, and except for a fever, no trace of infection was de-
tected by the ID physician. After a confirmation of signifi-
cant eosinophilia (2,173/µL), a negative blood culture, and
computed tomography showing negative results, piperacillin/
tazobactam administration was discontinued after 4 days. A
serum concentration of teicoplanin that remained (9.3 µg/
ML) for 7 days after the discontinuation may have been the
cause of the sustained fever.

Among most (91%, 10/11 cases) of our study subjects, the
PCT levels were either negative or low. PCT is a pro-
hormone of calcitonin and is known to be generated during
bacterial infections (16). However, PCT false-positive cases
have also been reported in cases of major surgery, graft-
versus-host disease, and autoimmune disorders. A definitive
report regarding the relationship between PCT and DF has
not yet been reported in the pertinent literature. Delèvaux et
al. analyzed the PCT levels in a bacterial/fungal infection
group and an abacterial inflammatory group (17). In their
study, the PCT levels of only 2 DF patients were available,
and both were negative (0.1 ng/mL) (17). As mentioned
above, the most critical and difficult differential diagnosis of
DF is bacteremia without a clear focus. Schuetz et al. sug-
gested a cut-off level of 0.25 ng/mL for excluding bac-
temia (18). However, we do not always recommend that doc-
tors measure the PCT level due to the inherent limitations of
biomarkers. Following a detailed physical examination after
the correct microbiological tests have been performed to di-
gnose febrile patients, PCT may then be helpful in diag-
nosing DF.

There are several limitations associated with this study.
First, because we performed a retrospective study, some
clinical findings may have been missed, although all cases
were confirmed by ID physicians. To resolve these prob-
lems, we will continue to accumulate the data of our pa-
patients for a follow-up prospective analysis. Second, because
our hospital is a tertiary care medical center, which admits
critical patients from other hospitals, two or more antibiotics
are often simultaneously administered in each patient. We
were unable to clarify the causal drug for the 6 DF patients.
Third, other antibiotics were administered to most of our pa-
patients (10/16 cases) after the previous drugs had been
stopped. This was because we could not discontinue antibi-
otics for primary infectious illnesses and could not rule out
the possibility that other new nosocomial infections had oc-
curred.

In conclusion, among 388 cases, 16 participants (4.1%) were
diagnosed with DF during ID consultation. Their clini-
cal characteristics were similar to those of previous reports.
Clinicians must consider the possibility that glycopeptides,
similar to β-lactams, may be the causative agents of DF. Ad-

### Table 3. Laboratory Tests of Patients with Drug
Fever.

<table>
<thead>
<tr>
<th>Test</th>
<th>Mean ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak WBC, mean ± SD (µL)</td>
<td>7,519 ± 3,551</td>
</tr>
<tr>
<td>(Eosinophils, mean ± SD)</td>
<td>(420 ± 575)</td>
</tr>
<tr>
<td>Peak AST level, mean ± SD (IU/L)</td>
<td>47.7 ± 46.0</td>
</tr>
<tr>
<td>Peak ALT level, mean ± SD (IU/L)</td>
<td>48.9 ± 59.5</td>
</tr>
<tr>
<td>Peak CRP level, mean ± SD (ng/dL)</td>
<td>5.1 ± 3.9</td>
</tr>
<tr>
<td>Peak PCT level a</td>
<td></td>
</tr>
<tr>
<td>≤ 0.1 (ng/dL)</td>
<td>8</td>
</tr>
<tr>
<td>0.1 - 0.25 (ng/dL)</td>
<td>2</td>
</tr>
<tr>
<td>&gt; 0.25 (ng/dL)</td>
<td>1</td>
</tr>
<tr>
<td>Neutropenia</td>
<td>4</td>
</tr>
<tr>
<td>Eosinophilia</td>
<td>4</td>
</tr>
</tbody>
</table>

a: Not available in five patients. AST: aspartate aminotransferase, ALT: alanine aminotransferase, CRP: C-reactive protein, PCT: procalcitonin, SD: standard deviation, WBC: white blood cell.
ditionally, a negative or low procalcitonin level may be suggestive of DF.

The authors state that they have no Conflict of Interest (COI).

Acknowledgement
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References