Management of febrile neutropenia in the United Kingdom: time for a national trial?

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Recent advances in febrile neutropenia (FN) have highlighted the value of risk stratification and the evolving role of oral antibiotics with early hospital discharge in low-risk patients. The aim of this study was to survey whether these advances have been translated into routine clinical practice in the UK. Questionnaires were sent to cancer clinicians across the UK to determine clinicians’ routine management of FN, including use of risk stratification, antibiotic regimen and criteria for hospital discharge. In all, 128 clinicians responded, representing 50 cancer departments (83%). Only 38% of respondents stratify patients according to risk and with substantial variation in the criteria defining ‘low-risk’. Furthermore, only 22% of clinicians use oral antibiotics as first-line treatment in any patients with FN, but this was significantly greater among clinicians who do compared to those who do not stratify patients by risk, 51 vs 4% (P<0.0001). These findings suggest a slow and/or cautious introduction of newer strategies for the management of low-risk FN in the UK. However, 84% of respondents confirmed their willingness to participate in a trial of oral antibiotics combined with early discharge in low-risk FN.

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For more than 30 years the standard management for all patients developing febrile episodes while neutropenic has been in-patient treatment with broad spectrum intravenous (i.v.) antibiotics (Hughes et al, 1998; Bodey et al, 1997; Hughes et al, 1997). However, recent evidence has prompted a reassessment of febrile neutropenia (FN), with increasing realisation that such intensive treatment may not be necessary or appropriate for all patients. This viewpoint has stemmed from the recognition that FN is not a single entity but rather represents a spectrum of severity. Only a small proportion of patients with FN will develop serious medical complications (16% overall, 12% of patients with solid tumours/lymphomas and 18% of patients with haematological malignancies) and fewer than 5% will die as a result of the episode (4.8%; Klastersky et al, 2000). Thus a majority of episodes of FN may be described as being ‘low-risk’. Several groups of investigators have independently developed prognostic indices in FN in an attempt to identify criteria by which to define ‘low-risk’ (Rubin et al, 1988; Talcott et al, 1992; Viscoli et al, 1994; Elting et al, 1997; Talcott et al, 1998). More recently these groups have come together in an international collaboration with the publication of the Multinational Association for Supportive Care in Cancer (MASCC) ‘risk index’ (Klastersky et al, 2000). In this study more than 40 possible risk factors for the development of significant medical complications at presentation of FN were examined in an initial derivation set of 756 patients. From this, logistic regression analysis was used to derive a model for the risk of development of such complications, which was subsequently tested in a further validation set of patients. The resulting risk index consists of seven weighted clinical factors (Table 1). The value of this scoring system in the prediction of the development of serious complications has subsequently been validated in both the single-centre (Uys et al, 2004) and multicentre settings (Paesmans et al, 2003).

The aim of defining low-risk neutropenia in this way has been to identify episodes of FN, which may be amenable to newer approaches to treatment involving less intensive, more convenient treatment, which may in turn be expected to have an impact on patients’ quality of life. Such treatment strategies may also have a potentially significant impact on health service resource utilisation.

Most recent research effort has focused on the role of oral antibiotics. Two large multinational prospective randomised controlled trials (Freifeld et al, 1999; Kern et al, 1999) have demonstrated equivalence in terms of both efficacy and safety for oral antibiotics compared to standard parenteral regimens for patients with low-risk FN when delivered in the in-patient setting. These findings have led to the inclusion of combination oral antibiotics as a standard option for treatment of low-risk FN in the most recent Infectious Diseases Society of America guidelines for the use of antimicrobial agents in neutropenic patients with cancer (Hughes et al, 2002). The case for oral antibiotics has been further strengthened by a recent meta-analysis (Vidal et al, 2004) of 15 trials comparing oral and i.v. treatment, which found no difference in failure rates or mortality between the two interventions.

The feasibility of combining an oral antibiotic regimen with early hospital discharge has been reported in a UK population in a randomised single-centre trial (Innes et al, 2003) which compared this approach to standard in-patient i.v. treatment. In this study,
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tionnaires to distribute to other relevant colleagues.
rationale, a stamped addressed envelope and additional ques-
September 2003 with a covering letter explaining its aims and
of FN. Questionnaires were sent to these identified clinicians in
across the UK with an interest in the antibiotic management
249 consultant oncologists and haematologists from 60 centres
mainly tick-box responses. The name and hospital of the clinician
asked to return the survey uncompleted, but otherwise to proceed
asked whether they manage patients with FN. If not they were
interested to determine:
(i) Whether UK clinicians are assessing the likelihood of patients
developing complications related to FN and if so, whether
they are using the MASCC index or other means.
(ii) Whether UK clinicians are using oral antibiotics as first-line
treatment in any patients with FN.
(iii) What criteria UK clinicians are using to determine patients’
suitability for hospital discharge and whether they have
policies for early hospital discharge.

METHODS
A database of clinicians registered to a UK national trial of
prophylactic antibiotics (Cullen et al, 2005) was used to identify
249 consultant oncologists and haematologists from 60 centres
across the UK with an interest in the antibiotic management
of FN. Questionnaires were sent to these identified clinicians in
September 2003 with a covering letter explaining its aims and
rational, a stamped addressed envelope and additional question-
naires to distribute to other relevant colleagues.

The questionnaire
The questionnaire was designed to be succinct (eight questions
covering one sheet of paper) and easy to answer consisting of
mainly tick-box responses. The name and hospital of the clinician
was preprinted onto the questionnaire. Clinicians were initially
asked whether they manage patients with FN. If not they were
asked to return the survey uncompleted, but otherwise to proceed
to answer the remaining questions. The questions related to
the three key areas specified below and clinicians were invited to also
give further comments.

Stratification and treatment of low-risk FN Clinicians were
asked whether they routinely stratify patients with FN and, if so,
to explain the criteria by which they defined ‘low-risk’. They were
then asked about their standard first-line antibiotic treatment for
non-penicillin-allergic low-risk patients with FN. They were asked
to specify whether they use an i.v. antibiotic regimen until
resolution, an i.v. regimen initially followed by ‘step-down’ oral
antibiotics or oral antibiotics from the outset and for the details of
the regimen used. There was also an additional question asking
about the timing of antibiotic discontinuation.

Criteria for hospital discharge Clinicians were asked for the
criteria they use for hospital discharge for patients recovering from
febrile neutropenia. They were asked whether they take into account
temperature, neutrophil count, a combination of both or other
criteria. If temperature was used as a criterion for discharge
clinicians were asked to further define the absolute value and
duration of the patient’s defervescence, which would be
deemed appropriate for discharge. Similarly, clinicians who
specified neutrophil count as a criterion were asked for the
absolute neutrophil count which would be deemed appropriate
for discharge or alternatively whether they used a rising neutrophil
count, irrespective of the absolute value as their preferred
criterion.

Trial participation Finally, clinicians were asked whether they
would be willing to participate in a trial for low-risk FN using oral
antibiotics in conjunction with early hospital discharge.

Statistical analysis
The responses of clinicians to the questionnaire are reported
descriptively. Type of antibiotic treatment and criteria for hospital
discharge are compared between those who do or do not stratify
patients by risk using continuity-adjusted χ² tests.

RESULTS
Of the 249 questionnaires sent out to clinicians from 60 cancer
departments across the UK, replies were received from 114 (47.4%)
clinicians who manage patients with FN. A further 14 replies were
received from clinicians who were not included in the original
mailshot but who had been passed copies of the questionnaire
by other recipients. Thus a total of 128 responding clinicians
(94 clinical/medical oncologists; 32 haematologists and two others)
who manage FN responded. These represent 50 departments
across the UK with a median of two clinicians from each centre
(rage 1–7).

Stratification of FN
Of these 128 clinicians, 79 (62%) do not stratify patients with FN
into low- and high-risk categories (Figure 1). Of the 49 (38%)
who do stratify, three indicated that this was on an occasional or
informal basis. Of the 50 oncology/haematology departments
represented, 28 (56%) had at least one clinician who stratifies
patients by risk and 22 (44%) had no responding clinician who
stratifies such episodes by risk. Within the centres represented
there was a lack of consistency regarding risk stratification;
of the 29 centres from which two or more clinicians responded,
there were only five centres from which all responding clinicians
stratify by risk.

Of the 49 clinicians who stratify by risk, 43 gave the criteria by
which they define ‘low-risk’, three stated that there were no formal
criteria and three gave no further information. The criteria varied
substantially, with the majority using a combination of factors
including patients’ symptoms and signs, absolute neutrophil count
and expected duration of neutropenia, nature of the underlying
malignancy, previous episodes of neutropenia and age (Table 2).
However, very few clinicians appeared to be systematically using
a single published definition of low-risk but rather to be using
factors derived from more than one study. In addition, some
clinicians gave more general and subjective definitions, for

Table 1 MASCC Risk index

| Characteristic | Score |
|----------------|-------|
| Burden of illness | 6 |
| No or mild symptoms* | 5 |
| Moderate symptoms* | 3 |
| No hypotension | 5 |
| No chronic obstructive pulmonary disease | 4 |
| Solid tumour/lymphoma or no previous fungal infection | 4 |
| No dehydration | 3 |
| Outpatient status at onset of fever | 2 |
| Age < 60 years | 1 |

*Points attributable to burden of illness are not cumulative. The maximum
theoretical score is therefore 26. The authors used a threshold of ≥21 points to
define low-risk.

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Figure 1

Summary of clinicians’ antibiotic treatment of low-risk FN.

Table 2

| Factor used to stratify risk of FN | No of clinicians using this factor |
|-----------------------------------|-----------------------------------|
| Clinical findings                 | 26                                |
| Absolute neutrophil count         | 17                                |
| Anticipated duration of neutropenia | 16                                |
| Patients’ symptoms or performance status | 12                                |
| Underlying malignancy (nonhaematological or nonacute leukaemia) | 9                                  |
| Site of infection                 | 8                                 |
| Comorbidities                     | 7                                 |
| Age                               | 5                                 |
| Chemotherapy regimen              | 4                                 |
| Previous fungal infection in haematological malignancy | 3                                  |
| Outpatient at presentation        | 2                                 |
| Previous episodes of febrile neutropenia | 1                                  |
| ‘Controlled’ cancer               | 1                                 |

Table 3

| Antibiotic regimen used | No. (% of known) clinicians using |
|-------------------------|----------------------------------|
| Oral antibiotic regimen (n = 28) |                                    |
| Ciprofloxacin+co-amoxiclav | 18 (69)                          |
| Ciprofloxacin            | 8 (31)                           |
| Not specified            | 2                                 |
| I.V. antibiotic regimen (n = 79) |                                    |
| Dual therapy             | 40 (82)                          |
| Monotherapy              | 9 (18)                           |
| Not specified            | 7                                 |
| Oral antibiotic regimen (n = 43) |                                    |
| Dual therapy             | 25 (68)                          |
| Monotherapy              | 12 (32)                          |
| Not specified            | 6                                 |

Criteria for hospital discharge

The criteria used by respondents for hospital discharge are shown in Table 4. The majority (n = 91; 71%) used a combination of temperature and neutrophil count as criteria for hospital discharge (including four who stated that just one criteria is used in certain situations and one whose criteria also included five days i.v. antibiotics). Of the remaining respondents, 33 (26%) used only the patient’s temperature, two (2%) used only the neutrophil count, one specified that discharge was at the clinician’s discretion and one did not specify. Those who stratified patients according to risk were more likely to utilise fever lysis alone, irrespective of neutrophil count: 17 of the 49 (35%) clinicians who do stratify by risk use temperature only compared with 16 of the 79 (20%) nonstratifying clinicians although this did not reach statistical significance (P = 0.11). Of the 123 clinicians who specified temperature criteria for discharge, almost half (n = 60; 49%) specified ≤37°C and half (n = 60; 49%) specified ≤37.5°C, with 24 h the most popular duration (n = 87; 71%). Of the 93 clinicians who gave patient’s neutrophil count as part of their criteria for discharge approximately a third (n = 32; 34%) use a rising neutrophil count irrespective of the absolute value, while the remainder (n = 61; 66%) have a fixed absolute neutrophil count which they regard as being suitable for discharge.

Duration of antibiotic treatment

The majority of clinicians (n = 79; 62%) stated that in low-risk patients antibiotics were discontinued after a minimum number days following lysis of fever with 1–2 days specified by 14 clinicians, 2–5 days by 54 and 5–7 days by 11. A further 27 (21%) discontinue antibiotics at (or 24 h before; n = 1) discharge. Nine clinicians (7%) discontinue after a fixed duration from onset of fever with number of days ranging from 3 to 10. The criteria for the remaining clinicians related to neutrophil recovery (n = 4) or white blood count (n = 1) and eight not specifying.

Clinicians’ comments and trial participation

When asked for their further comments, clinicians from three centres commented that they manage some patients with low-risk FN as outpatients and a further five that they have early discharge policies in place. A number of clinicians also made comments regarding practical difficulties in the assessment and management of low-risk FN. In particular, several respondents cited difficulties with the ability of junior staff to assess risk, for example ‘junior staff play it safe’; ‘difficulty in determining whether) the nadir is passed or coming’. One clinician commented that ‘in practice very
rarely do patients get into low-risk category’. However, respondents were generally very favourable about the concept of early discharge for patients with low-risk FN, one clinician commenting that ‘we have real pressure on beds, anything reducing in-patient stay would be of interest’. Of the 128 respondents, 108 (84%) said that they would be willing to participate in a trial for low-risk FN of oral antibiotics combined with early hospital discharge.

DISCUSSION

This study is the first to survey the UK management of FN, an important complication of anticancer chemotherapy. Clinicians from 50 of the 60 cancer departments surveyed (83%) responded, giving broad representation of current treatment across the UK. The findings suggest substantial variation in the management of FN and a slow and/or cautious introduction of recent evidence in this area.

Fewer than half of the responding clinicians stratify patients with FN according to their likelihood of developing complications. Moreover, of those who do stratify, the definition of ‘low-risk’ varies greatly, with clinicians using different combinations of a variety of published and unpublished, objective and subjective criteria. Clinicians from only one centre (the author’s) were using the well-validated MASCC scoring system. Moreover, only a quarter (22%) of respondents use oral antibiotics from the outset in the management of any of their patients with FN. This is despite the increasing evidence base, which demonstrates equivalence in terms of success rates and safety for oral vs i.v. regimens in low-risk FN (Freifeld et al, 1999; Kern et al, 1999; Innes et al, 2003), the anticipated benefits of quality of life to patients and the potential significant savings in utilisation of resources. Furthermore, the study findings represent the practice of clinical trialists in the UK in the field of FN. It may be expected that as such they are more likely to have incorporated recent study findings into their clinical practice. Therefore, this study may overestimate the use of oral antibiotics in the routine management of FN in the UK. The reasons for not implementing such policies have not been explored in the current survey, but it is possible that some clinicians feel that the level of staffing available is inadequate to safely implement the close monitoring required for such policies.

Interestingly, clinicians and centres that do stratify and thus identify low-risk patients were more likely to have introduced oral antibiotics into their management of FN than those who do not stratify. Thus acknowledgement of the concept of ‘low-risk’ is accompanied by recognition that a ‘one size fits all’ management strategy is not necessarily the most appropriate treatment of FN. Similarly, clinicians who stratified by risk were more likely to use only patients’ temperature rather than temperature together with neutrophil count as criteria for determining discharge from hospital than those who do not stratify by risk. In turn this is likely to reflect shorter durations of hospital admission for stratifying clinicians/centres, although this has not been formally examined.

The published evidence presents a strong case that the introduction of less intensive management strategies could offer significant benefits to patients with low-risk FN in terms of reduced hospitalisation and potentially improved quality of life. In a previous UK study, it was estimated that the approximate costs per episode of low-risk FN treated with oral antibiotics/early discharge were 55% of those for a patient treated as an in-patient treated with i.v. antibiotics (Innes et al, 2003). Approximately 70% of all FN episodes are low-risk (Klastersky et al, 2000). Thus, if such a management strategy could be safely achieved across the UK, the potential resource savings would be considerable.

There is more than one strategy for using the limited available repertoire of orally bioavailable antibacterials to combat neutropenic infection. Instead of using fluoroquinolones to treat neutropenic infections for patients predicted to be at low risk of complications, they might be used at an earlier time point as prophylaxis. Recent results from a randomised, double-blind, placebo-controlled trial of 1565 patients with solid tumours or lymphoma (Cullen et al, 2005) have shown reduction in the frequency of febrile episodes during chemotherapy in patients receiving levofloxacin prophylaxis compared with those receiving placebo (11 vs 16%; P = 0.01). These important results are not in themselves sufficient to recommend a universal prophylaxis strategy because the impact on bacterial resistance patterns has not yet been determined. A targeted treatment approach for actual infections using oral fluoroquinolones might enable less intensive treatment and early hospital discharge and these potential benefits need to be assessed. It is not known whether patients who develop FN despite receiving prophylactic antibiotics could be still be defined as ‘low-risk’ even if they otherwise fulfil the MASCC criteria, since such patients have been excluded from previous studies. Further research in this area is required.

In conclusion, the present study identifies both variations in practice and a degree of caution in implementing evidence-based advances in the management of FN in the UK. Better local and national policies and guidelines are clearly needed. We believe that

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Table 4  Criteria used for patient discharge. (a) Temperature and duration for those who use temperature criterion and (b) Neutrophil count criteria for those who use it

| Criteria | 1 reading | 2 readings 4 h apart | 3 readings 4 h apart | 24 h | Other* | Total |
|----------|-----------|----------------------|---------------------|------|--------|-------|
| (a)      |           |                      |                     |      |        |       |
| ≤ 37°C   | 0         | 8                    | 1                   | 44   | 7      | 60    |
| ≤ 37.5°C | 2         | 10                   | 3                   | 4    | 4      | 60    |
| ≤ 37.9°C | 0         | 1                    | 0                   | 2    | 0      | 3     |
| Total    | 2         | 19                   | 4                   | 87   | 11     | 123   |

| Criteria | 1 reading | 2 readings 4 h apart | 3 readings 4 h apart | 24 h | Other* | Total |
|----------|-----------|----------------------|---------------------|------|--------|-------|
| (b)      |           |                      |                     |      |        |       |
| Neutrophil count only | 0 | 36 (and rising in 1) | 0                   | 36   | (39%) |     |
| ≥ 0.5 x 10^9/l | 1 | 18 (and rising in 3) | 3                   | 22   | (24%) |     |
| ≥ 1 x 10^9/l | 1 | 29                   | 2                   | 32   | (34%) |     |
| Rising neutrophil count irrespective of value | 1 | 3                   | 0                   | 3    | (3%)  |     |
| Other*  | 0         | 3                    | 0                   | 5    | 93     |      |

*Includes 72 h, 1 reading or 24, 24–48, 48, 72, > 24 h.
*One respondent who stated temperature and neutrophil count as criteria for patient discharge did not actually specify the criteria.
*Includes 0.8 × 10^9/l, 0.3 × 10^9/l, 0.2 × 10^9/l and rising.
*Two of the clinicians who specified ‘other’ criteria did not use neutrophil count.
the introduction of newer strategies may be best achieved in the setting of a well-conducted, multicentre trial. The advent of National Cancer Research Networks across the UK provides a unique opportunity for such a study. A randomised phase III trial comparing standard in-patient management with early hospital discharge in cancer patients receiving oral antibiotics for low-risk FN has been sponsored by Cancer Research UK and will be launched later in the year.

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