It is feasible to screen for important RA co-morbidities using patient completed questionnaires

Background: To study the feasibility of pre-consultation questionnaires to screen for Rheumatoid Arthritis (RA) co-morbidities compared with face-to-face consultation. Methods and findings: Postal questionnaires were sent to all RA patients on our electronic database. Four domains were studied: Gastrointestinal (GI) symptoms, Cardiovascular (CVS) symptoms and risk factors; Foot; Anxiety and depression (Hospital Anxiety and Depression Scores (HADS)). Patients were then reviewed clinically, with a full history and examination undertaken and HAQ, DAS and QRISK2 scores were calculated for each patient. 597 questionnaires were sent, 301 (50%) completed and 129 patients were reviewed in clinic. The mean DAS score was 3.04. 32% of responders reported daily GI symptoms, with no extra symptoms of concern elicited at clinic review in any patient. 49% patients had high CVS risk as calculated by QRISK2. 5% patients assessed as low CVS risk by our questionnaire were re-categorised as high risk by QRISK2. 89% of patients reported foot symptoms in the past week. Mean foot score was 4/10 with correlation co-efficient 0.63 between postal and clinical scores. The mean anxiety score was 7.8 with 49% patients scoring 8 or more. The mean depression score was 9.3 with 64% patients scoring 8 or more. Conclusion: Pre clinic screening by questionnaire is a feasible way of identifying comorbidities. We found a high rate of co-morbidities in our RA population.

Keywords: rheumatoid arthritis • co-morbidities • health screening • questionnaires
the tender joint count and VAS components of
the Disease Activity Score (DAS).

Aims
We assessed the feasibility of using postal
questionnaires to assess comorbidities by
comparing results obtained from postal
questionnaire with face to face consultation. We
also documented the prevalence of extra-articular
symptoms and comorbidities associated with RA
in our patient population.

Methods
Postal questionnaire
A postal questionnaire was sent to all patients
with a diagnosis of RA, confirmed by a
consultant rheumatologist, on our electronic
drug monitoring database (Supplementary
File). They were also invited to participate in a
general health screening research clinic and were
asked to return a signed consent form with the
questionnaire if they wished to attend this.
The questionnaire was divided into 4 sections:

1. Section 1 asked 10 questions regarding
   GI health:
   Patients were asked to indicate the frequency of
   10 GI symptoms with a maximum score of 30.

2. Section 2 assessed 8 CVS symptoms and
   risk factors:
   The maximum CVS score was 10. Total CVS
   scores of 2 and 3 were compared as thresholds to
   indicate an increased risk of cardiovascular event.
   Patients were asked about type and frequency of
   exercise.

3. Section 3 assessed foot symptoms using
   the 10 point Swindon Foot and Ankle
   Questionnaire (SFAQ):
   A foot score out of 10 was assigned; with
   the answer no scoring 0 and yes scoring 1
   (Supplementary File for more details on scoring).

4. Section 4 contained a Hospital Anxiety
   and Depression (HAD) questionnaire
   [9,10].
   HAD anxiety and depression scores were
   calculated with a maximum score of 21. The cut
   off of 8 or more was used to indicate patients at
   risk of anxiety or depression [11].

Readability
The questionnaires were designed to have
a reading age of 11. This was to improve
collection rates by patients, because 16%
of adults in the UK are functionally illiterate,
defined as a reading age at or below that of an 11
year old [12-14].

Face to face consultation
Patients underwent a comprehensive clinical
review including:

1. Full GI history and examination, including
   for periodontal and perianal
disease. GI score out of 30 was
   recalculated after clinical assessment.

2. QRISK2 score calculation (with blood
   pressure, blood sugar and cholesterol
   checked) along with a CVS examination
   [15]. A QRISK2 score of >20% CVS
   event risk at 10 years was considered as
   high risk.

3. Detailed foot examination (including
   completion of the validated Manchester
   Foot Pain and Disability Index (MFPDI)
   [16,17].

4. HAD was reassessed.

5. Activity of RA was documented using
   a Disease Activity Score (DAS) and
   an Overall Status in Rheumatoid
   Arthritis (OSRA) score [18,19]. Patients
   also completed a Health Assessment
   Questionnaire (HAQ), a Euroqol
   score was calculated and a medication
   history was taken, including use of
   complementary medications and
   therapies [20-22].

6. Demographic data was collected,
   including employment status.

Ethical approval was obtained from Wiltshire
Research Ethic Committee and written consent
obtained from all patients. Microsoft Excell
2007 was used to perform statistical analysis.
All correlation co-efficients were calculated as
Spearman rank correlation coefficients.

Results
Of 597 questionnaires sent, 301 (50.3%) were
returned. 129 patients (43% of questionnaire
responders and 22% of total RA population sent
screening questionnaires) were seen in research
clinic.

Of the 301 patients returning questionnaires
202 (67%) were female, with an age range of 31–
91 (median 62). Of the 129 patients reviewed
in clinic, 90 (61%) were female, with an age
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Gastrointestinal health

32% of questionnaire responders and 18% of patients seen in clinic reported at least 1 daily GI symptom (Table 1). 17% and 9% of patients respectively reported no GI problems. There were no statistically significant differences in frequency of GI symptoms reported on questionnaire compared with clinic. Out of the symptoms reported on at least a weekly frequency, indigestion, bloating and abdominal pain were the most commonly reported (Figures 1A and 1B).

The only significant difference in specific symptom reporting between the questionnaire and clinic was steatorrhoea, with no patients experiencing true steatorrhoea on questioning in clinic but 4% reporting it daily, 8% weekly and 13% monthly on questionnaire.

Extra information obtained at clinic review:

1. Dental disease:
   - 109 (84%) had fillings;
   - 11 (<1%) patients had normal dentition;
   - 28 (22%) had poor gum health, including loose teeth or gingivitis;
   - 70 (54%) patients had a dry or coated tongue;
   - 42 (32%) of patients reported mouth ulcers.

2. 36 patients (28%) had haemorrhoids.

3. Use of medication with commonly reported GI side effects (GI toxic drugs):

The mean total GI score of all 301 questionnaires was 5 (range 0–21). The mean score at clinic review was 4 (range 0–12). Of the patients seen in clinic, the mean GI score from their questionnaires was 5 (range 0–19). The correlation coefficient of their clinic score compared with their questionnaire score in those patients who were assessed in clinic was 0.64 (Figure 2).

<table>
<thead>
<tr>
<th>Symptom frequency</th>
<th>Number of symptoms</th>
<th>1 or more</th>
<th>2 or more</th>
<th>3 or more</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Questionnaire 301 pts</td>
<td>Clinic 129 pts</td>
<td>Questionnaire 301 pts</td>
<td>Clinic 129 pts</td>
</tr>
<tr>
<td>Daily</td>
<td>96 (32%)</td>
<td>23 (18%)</td>
<td>35 (12%)</td>
<td>5 (4%)</td>
</tr>
<tr>
<td>At least weekly</td>
<td>160 (53%)</td>
<td>68 (53%)</td>
<td>96 (32%)</td>
<td>39 (30%)</td>
</tr>
<tr>
<td>At least monthly</td>
<td>246 (82%)</td>
<td>119 (92%)</td>
<td>205 (68%)</td>
<td>85 (66%)</td>
</tr>
<tr>
<td>Never</td>
<td>50 (17%)</td>
<td>11 (9%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Frequency of GI symptoms.

Figure 1A. Frequency of GI symptoms from postal questionnaire.
• 34 (26%) of patients were on bisphosphonates;
• 12 patients (9%) were on prednisolone;
• 64 patients (50%) were on NSAIDS;
• 15 patients (12%) were on at least 2 of the above.

Patients taking potentially GI toxic drugs were more likely to suffer GI symptoms on at least weekly basis—49% vs. 34% (Table 2).

4. 52 patients (40%) were on a Proton Pump Inhibitor (PPI). 31/52 (60%) were taking at least 1 GI toxic drugs. Co-prescription of a PPI did not reduce the reporting rate of GI symptoms, whereas those patients taking a PPI but no GI toxic medication did have a lower GI symptoms rate.

Cardiovascular disease screening

121 patients had a both a postal score and Qrisk® 2 score in clinic calculated. The mean and median postal score was 2 (range 0- 6). Using 2/10 postal score as the threshold for high CVS risk, 85 (70%) patients were deemed high risk. If 3/10 postal score was used, 56 (46%) patients
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were deemed high risk. 51 (41%) patients were stratified as high risk by QRisk’2.

76 patients (63%) were classified in the same risk group using a postal cut off of 2 or more compared with QRisk’2 score. 77 patients (64%) were classified in the same risk group using a postal cut of 3 or more (Table 3).

5 patients (4%) deemed low risk on postal questionnaire using a cut off of 2 were reclassified as high risk by QRisk’2 (false negatives). 22 patients (18%) patients using a cut off of 3 were deemed low risk by postal questionnaire but were high risk on QRisk’2. 40 patients (33%) deemed high risk on postal questionnaire using a cut off of 2 were reclassified as low risk by QRisk’2 (false positive). 22 patients (18%) patients using a cut off of 3 were deemed low risk by postal questionnaire but were high risk on QRisk’2 score.

284 patients answered the questions about exercise on the postal questionnaire. Of these, 86 patients (30%) never exercised, 21 (7%) exercised monthly, 75 (26%) weekly and 103 (36%) daily. 14 patients did both cardiovascular and stretching exercises. 76 performed stretching only, 42 performed cardiovascular only, 67 used walking and 4 used gardening as their main exercise.

7 patients were found to have previously undiagnosed hypertension. 32 patients (26%) were found to have cholesterol of greater than 4. 42 patients (35%) were taking a statin.

12 patients were taking oral steroids and 64 patients were taking NSAIDs. 23 of these patients (32%) had a high QRisk’2 score compared with 16 patients (28%) not taking these medications, with a relative risk of 1.14.

Feet

From 301 questionnaires returned the mean Swindon Foot and Ankle Questionnaire Score (SFAQ) was 4 (range 0–10). 16 patients (11%) scored 0. The mean SFAQ foot score from 139 patients seen in clinic was 4 (range 0–10). The correlation between postal score and clinic score was r=0.63. The correlation between clinic score and the Manchester Foot score was 0.67 (Figure 3).

92% of patients assessed in clinic had some form of foot or ankle problems documented and 66% had more than one problem (Table 4).

Anxiety and depression

From 296 postal questionnaires, the mean anxiety score was 7.8 (range 0–21), with 146 patients (49%) scoring 8 or more. From 296 postal questionnaires, the mean depression score was 9.3 (range 0–21), with 188 patients (64%) scoring 8 or more. The anxiety and depression scores correlated with each other, with R=0.76 but did not correlate with other measures of disease outcome, including HAQ, DAS, GI score, foot score or exercise (Table 5A).

DAS28

Mean DAS score was 3.04 (range 1.1–6.19) and mean HAQ score was 1.25 (range 0–3).

The DAS28 did not correlate with any of the other domains assessed. The correlation coefficient for DAS28 and SFAQ was 0.29 (Table 5B).

Discussion

High rate of comorbidities once again demonstrated

We present the results of a screening exercise in a

<table>
<thead>
<tr>
<th>Table 2. GI symptoms and PPIs.</th>
<th>GI toxic med</th>
<th>No GI toxic meds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weekly GI symptoms</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ PPI</td>
<td>37/75 (49%)</td>
<td>18/53 (34%)</td>
</tr>
<tr>
<td>Weekly GI symptoms</td>
<td>16/31 (52%)</td>
<td>21/44 (47%)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 3. Comparison of CVS scores.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Risk category</td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>High</td>
</tr>
<tr>
<td>Low</td>
</tr>
<tr>
<td>Both scores high</td>
</tr>
<tr>
<td>Both scores low</td>
</tr>
<tr>
<td>Postal high, QRISK2 low</td>
</tr>
<tr>
<td>Postal low, QRISK2 high</td>
</tr>
</tbody>
</table>
Figure 3. Correlation between postal + clinic scores.

Table 4. Prevalence of foot pathologies.

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Number of patients</th>
<th>% patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hallux valgus</td>
<td>56</td>
<td>41</td>
</tr>
<tr>
<td>Callous</td>
<td>27</td>
<td>20</td>
</tr>
<tr>
<td>MTP synovitis</td>
<td>25</td>
<td>18</td>
</tr>
<tr>
<td>Ankle varus</td>
<td>22</td>
<td>16</td>
</tr>
<tr>
<td>Claw toes</td>
<td>21</td>
<td>15</td>
</tr>
<tr>
<td>Moretons neuroma</td>
<td>20</td>
<td>15</td>
</tr>
<tr>
<td>Posterior Tibial tendonitis</td>
<td>16</td>
<td>12</td>
</tr>
<tr>
<td>Peroneal tendonitis</td>
<td>16</td>
<td>12</td>
</tr>
<tr>
<td>Hind foot pathology</td>
<td>15</td>
<td>11</td>
</tr>
<tr>
<td>Ankle pathology</td>
<td>14</td>
<td>10</td>
</tr>
<tr>
<td>Hammer toes</td>
<td>14</td>
<td>10</td>
</tr>
<tr>
<td>Other</td>
<td>10</td>
<td>7</td>
</tr>
<tr>
<td>Previous surgery</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>Cross over</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Achilles tendonitis</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Plantar fascitis</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>

Number of diagnoses(%)

<table>
<thead>
<tr>
<th>Number of diagnoses</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
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<tbody>
<tr>
<td>12</td>
<td>9</td>
<td>35</td>
<td>58</td>
<td>21</td>
<td>10</td>
<td>2</td>
</tr>
</tbody>
</table>

Table 5. (A and B) HADS correlations.

(A)

<table>
<thead>
<tr>
<th></th>
<th>Anxiety</th>
<th>Depression</th>
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</thead>
<tbody>
<tr>
<td>HAQ</td>
<td>0.28</td>
<td>0.38</td>
</tr>
<tr>
<td>DAS</td>
<td>0.14</td>
<td>0.03</td>
</tr>
<tr>
<td>Gl score</td>
<td>0.18</td>
<td>0.16</td>
</tr>
<tr>
<td>Foot score</td>
<td>0.02</td>
<td>0.05</td>
</tr>
<tr>
<td>Exercise</td>
<td>0.13</td>
<td>0.16</td>
</tr>
</tbody>
</table>

(B)

<table>
<thead>
<tr>
<th></th>
<th>Correlation co-efficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>HAQ</td>
<td>0.09</td>
</tr>
<tr>
<td>Foot score</td>
<td>0.29</td>
</tr>
<tr>
<td>HAD anxiety</td>
<td>0.14</td>
</tr>
<tr>
<td>HAD depression</td>
<td>0.03</td>
</tr>
<tr>
<td>Gl score</td>
<td>0.1</td>
</tr>
<tr>
<td>QRISK2 score</td>
<td>-0.02</td>
</tr>
</tbody>
</table>
group of patients with RA, whose demographics are compatible with a typical DGH population. 301 patients completed the postal questionnaire and 129 patients were seen in clinic. The results demonstrate that our patients with RA have a high rate of self-reported extra-articular diseases and symptoms, in all the domains we studied. Nearly one third of patients reported daily GI symptoms. Nearly half of our patients had a 10 year cardiovascular event risk of greater than 20%. 90% of patients reported some form of foot pathology or symptom in the past week, consistent with results of other studies [23-29]. Similarly half of our patients had an anxiety or depression score indicating that they were at risk of problems. This compares with the findings of recent meta-analysis of RA where 34% of patients had a HAD score greater than 8 and 15% greater than 11 [30]. Our results highlight the importance of assessing the global health of our RA patients, particularly in light of the well-established link between RA and increased risk of cardiovascular disease and the on-going excess mortality associated with RA despite treatment advances.

Correlation between postal and clinic results
The correlation between postal results and clinic results were moderate in all domains, suggesting that self-completed questionnaires completed prior to a clinic appointment, either by post sent out with an appointment letter or completed in the waiting room on the day of appointment could be a useful means of highlighting to the assessing physician the health concerns of the patient, which could help to guide and focus the subsequent consultation. Few new symptoms or problems of concern were elicited at clinic that was missed by our questionnaires. Simple screening tools are feasible
Currently there are no freely available tools to simultaneously assess several facets of health in patients with RA. There are several patient reported outcome measures, each covering separate domains both specifically for use in RA and in the wider medical population [31]. ‘Questionnaire fatigue’ is a real problem for both patients and clinicians because of the length of some assessment tools and the problems in interpreting questionnaire results.

Our postal questionnaire was designed to be simple and brief to both complete and interpret. It is important that any screening tools take account of the average UK reading age of 11, are brief enough to maximise full completion, whilst covering all domains. By balancing all of these factors, the maximum amount of valid information can be obtained. There is very limited published data on the readability of currently available medical questionnaires and none regarding tools specifically used in the assessment of RA. To our knowledge, our postal questionnaire is the first assessment tool used in RA to consider the readability of the tool and to document the reading age results.

Tensions and barriers to multisystem review
An early treat to target approach in RA has improved disease outcomes. It is important however, that the focus upon achieving a threshold DAS does not distract completely from the assessment of the multisystem effects and comorbidities associated with RA. In particular there is a growing concern that feet are being considered comorbidity rather than an integral feature of RA by use of the DAS28. A recent survey found that 50% of clinicians did not examine feet as they were not included in DAS [32]. Time constraints, rising patient referral number s and the pressure to adhere to new to follow up ratios are all factors leading to a perceived difficulty in ensuring holistic care.

There is tension between the overlapping roles of primary and secondary care. A recent multi-national cross-sectional study found that rates of detecting co-morbidities were variable [31]. The Arthritis Research UK Adult Inflammatory Arthritis Clinical Studies Group looking at co-morbidities discussed the current status of care in the UK and how best to address the gap between the standards expected by national guidance bodies and current practice [33]. They concluded that patient reported outcomes were as important as data collected by hospital staff. Current tools specifically developed to look at co-morbidities in RA focus mainly upon detecting risk factors of excess mortality rather than factors affecting morbidity as well.

There was no correlation between the scores in different domains. In particular, a high anxiety and/or depression score did not predict a high score in other domains as an explanation for high levels of self-reported symptoms. This suggests that problems in one area of global health were independent of another. There was also no correlation between high scores in the extra-articular domains we examined and the severity of RA as assessed by DAS score. This was true even of those patients with an elevated
anxiety and depression score. This is in contrast to other studies which have demonstrated a relationship between high self-reported anxiety and depression and VAS and tender joint counts [34]. The lower weightings given to these components of the DAS score may account for the lack of association seen in our study. It highlights that anxiety and depression are not merely a reaction to active inflammation. 

Study limitations
Our return rate of questionnaires was 50%, which is similar to the response rate of other postal surveys. It is possible that there was a selection bias, in that those patients returning questionnaires and consenting to a clinic review were more likely to report symptoms. There was however, no statistically significant difference in scores comparing those patients seen in clinic, with those patients who returned questionnaires only [35].

We have not been able to cover every domain of extra-articular disease in RA and sleep, fatigue and joint stiffness are important omissions from our study. There was also limited patient involvement in the development of the questionnaire, both in terms of question design.

These results are from a single centre at one time point in 2010 and so caution should be given in more widely interpreting results. However, 6 years after our data collection, assessment of extra-articular features and co-morbidities in RA remains relevant. The rate of comorbidities detected in our studies is comparable with other published data.

Conclusion
Pre clinic screening by questionnaire is an efficient, effective and feasible way to identify co-morbidities and direct treatment. Important symptoms and red flags were generally not missed by this approach.

This study highlights both the high rate of significant co-morbidities associated with RA, and the multi-system nature of the disease. We found clinically acceptable correlation between postal questionnaires results and the clinic findings.

This simple practical working model can be used prior to clinic appointments to allow clinicians to cover a comprehensive range of RA co-morbidities plus focus consultations towards important patient concerns. We anticipate that these questionnaires will be adapted to an electronic format for use in the waiting room prior to consultations.

References
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