Introduction

The American College of Rheumatology (ACR) most recently published recommendations for the use of disease-modifying antirheumatic drugs (DMARDs) and biologic agents in the treatment of rheumatoid arthritis (RA) in 2008 (1). These recommendations covered indications for use, monitoring of side effects, assessment of the clinical response to DMARDs and biologic agents, screening for tuberculosis (TB), and assessment of the roles of cost and benefit. The views expressed in this article are those of the authors and do not necessarily reflect the position or policy of the Department of Veterans Affairs or the United States government.

Guidelines and recommendations developed and/or endorsed by the American College of Rheumatology (ACR) are intended to provide guidance for particular patterns of practice and not to dictate the care of a particular patient. The ACR considers adherence to these guidelines and recommendations to be voluntary, with the ultimate determination regarding their application to be made by the physician in light of each patient’s individual circumstances. Guidelines and recommendations are intended to promote beneficial or desirable outcomes but cannot guarantee any specific outcome. Guidelines and recommendations developed or endorsed by the ACR are subject to periodic revision as warranted by the evolution of medical knowledge, technology, and practice.

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Significance & Innovations

- These 2012 recommendations update the 2008 American College of Rheumatology recommendations for the treatment of rheumatoid arthritis (RA).

- The recommendations cover the use of disease-modifying antirheumatic drugs and biologic agents in patients with RA, including switching between drugs.

- We address screening for tuberculosis reactivation, immunization, and treatment of RA patients with hepatitis, congestive heart failure, and/or malignancy in these recommendations, given their importance in RA patients receiving or starting biologic agents.

Dr. Singh has received consultant fees, speaking fees, and/or honoraria (less than $10,000 each) from Allergan, Ardea, Savient, and Novartis, and (more than $10,000) from Takeda, has received an investigator-initiated grant from Savient and Takeda, and is an executive member of an international organization, Outcome Measures in Rheumatology (OMERACT). Dr. Furst has received consultant fees, speaking fees, and/or honoraria (less than $10,000) from Abbott, Actelion, Amgen, BMS, Biogen Idec, UCB, Gilead, Centocor, GSK, Novartis, Pfizer, NIH, and Roche/Genentech, and is a member of the Consortium of Rheumatology Researchers of North America (CORRONA). Dr. Curtis has received consultant fees, speaking fees, and/or honoraria (less than $10,000 each) from Pfizer, BMS, Crescendo, and Abbott, and (more than $10,000 each) from Roche, Genentech, UCB, Centocor, CORRONA, and Amgen. Dr. Kavanaugh has conducted clinical research for Centocor, UCB, Genentech/Genentech, NIH, Abbott, Takeda, and Amgen. Dr. Kremer has received consultant fees, speaking fees, and/or honoraria (less than $10,000 each) from Pfizer, BMS, Crescendo, and Abbott, and (more than $10,000) from Roche, Genentech, UCB, Centocor, and Amgen. Dr. Moreland has received consultant fees (less than $10,000) from Pfizer and is a member of the data safety monitoring board for ChemoCentryx. Dr. Winthrop has received consultant fees, speaking fees, and/or honoraria (less than $10,000 each) from Abbott, Amgen, Pfizer, Celleristis, and Wyeth. Dr. Chatham has served as a paid consultant with investment analysts on behalf of Gerson Lehrman and Leerink-Swann. Dr. Bombardier has received honoraria (less than $10,000 each) and/or served on the advisory board for AbbVie Canada, AstraZeneca, Biogen Idec, BMS, Pfizer (Wyeth), Merck (Schering), Janssen (Merck), and Takeda, and has received honoraria (more than $10,000 each) from AbbVie International and Pfizer. Dr. Douchados has received consultant fees, speaking fees, and/or honoraria (less than $10,000 each) from Pfizer, Abbott, UCB, BMS, and Roche. Dr. Khanna has received consultant fees, speaking fees, and/or honoraria (less than $10,000 each) from AbbVie International and Pfizer, Actelion, and Gilead. Ms Leong has received consultant fees, speaking fees, and/or honoraria (more than $10,000 each) from Centocor Ortho Biotech and GlaxoSmithKline. Dr. Kolba owns stock and/or stock options in Merck, Amgen, and Genentech. Dr. Saag has received consultant fees, speaking fees, and/or honoraria (less than $10,000 each) from Merck, Lilly, Novartis, Genentech, Horizon, and URL, and (more than $10,000) from Amgen.

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the scenarios created using an ordinal scale specified in the RAND/University of California at Los Angeles (RAND/UCLA) Appropriateness Method (2–4). This method solicited formal input from this multidisciplinary TFP to make recommendations informed by the evidence. The methods used to develop the updated ACR recommendations are described briefly below.

**Systematic literature review: sources, databases, and domains.** Literature searches for both DMARDs and biologic agents relied predominantly on PubMed searches with medical subject headings and relevant keywords similar to those used for the 2008 ACR RA recommendations (see Supplementary Appendices 1 and 2, available in the online version of this article at http://onlineibrary.wiley.com/journal/10.1002/(ISSN)2151-4658). We included randomized controlled trials (RCTs), controlled clinical trials, quasi-experimental designs, cohort studies (prospective or retrospective), and case–control studies, with no restrictions on sample size. More details about inclusion criteria are listed below and in Supplementary Appendix 3 (available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658).

The 2008 recommendations were based on a literature search that ended on February 14, 2007. The literature search end date for the 2012 update was February 26, 2010 for the efficacy and safety studies and September 22, 2010 for additional qualitative reviews related to TB screening, immunization, and hepatitis (similar to the 2008 methodology). Studies published subsequent to that date were not included.

For biologic agents, we also reviewed the Cochrane systematic reviews and overviews (published and in press) in the Cochrane Database of Systematic Reviews to identify additional studies (3–8) and further supplemented by hand checking the bibliographies of all included articles.

### Table 1. Overview comparison of topics and medications included in the 2008 and 2012 American College of Rheumatology rheumatoid arthritis recommendations*

<table>
<thead>
<tr>
<th>Topic area considered</th>
<th>2008</th>
<th>2012</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indications for starting or resuming DMARDs and biologic agents</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>DMARDs included†</td>
<td>1. Hydroxychloroquine</td>
<td>1. Hydroxychloroquine</td>
</tr>
<tr>
<td>2. Leflunomide</td>
<td>2. Leflunomide</td>
<td></td>
</tr>
<tr>
<td>5. Sulfasalazine</td>
<td>5. Sulfasalazine</td>
<td></td>
</tr>
<tr>
<td>And, when appropriate, combination DMARD therapy with 2 or 3 DMARDs†</td>
<td>And, when appropriate, combination DMARD therapy with 2 or 3 DMARDs†</td>
<td></td>
</tr>
<tr>
<td>Biologic agents included§</td>
<td>Non-TNF</td>
<td>Non-TNF</td>
</tr>
<tr>
<td>1. Abatacept</td>
<td>1. Abatacept</td>
<td></td>
</tr>
<tr>
<td>2. Rituximab</td>
<td>2. Rituximab</td>
<td></td>
</tr>
<tr>
<td>Anti-TNF</td>
<td>3. Tocilizumab</td>
<td></td>
</tr>
<tr>
<td>3. Adalimumab</td>
<td>Anti-TNF</td>
<td></td>
</tr>
<tr>
<td>4. Etanercept</td>
<td>4. Adalimumab</td>
<td></td>
</tr>
<tr>
<td>5. Infliximab</td>
<td>5. Etanercept</td>
<td></td>
</tr>
<tr>
<td>Role of cost and patient preference in decision making for biologic agents</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Switching between therapies</td>
<td>Considered, but not addressed in detail</td>
<td>See 2008 recommendations¶</td>
</tr>
<tr>
<td>Monitoring of side effects of DMARDs and biologic agents</td>
<td>✓</td>
<td>See 2008 recommendations¶</td>
</tr>
<tr>
<td>TB screening for patients starting/receiving biologic agents</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Use of biologic agents in high-risk patients (those with hepatitis, congestive heart failure, and malignancy)</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Vaccinations in patients starting/receiving DMARDs or biologic agents</td>
<td>Pneumococcal, influenza, and hepatitis vaccines</td>
<td>Pneumococcal, influenza, hepatitis, human papillomavirus, and herpes zoster vaccines</td>
</tr>
</tbody>
</table>

* DMARDs = disease-modifying antirheumatic drugs; non-TNF = non–tumor necrosis factor; TB = tuberculosis.
† Cyclosporine, azathioprine, and gold were included in the literature search, but due to the lack of new data and/or infrequent use, they were not included in scenarios and the recommendations.
‡ Triple therapy with methotrexate + hydroxychloroquine + sulfasalazine.
§ Anakinra was included in the literature search, but due to the lack of new data and/or infrequent use, it was not included in the recommendations.
¶ No significant new data related to these topics.
Finally, the CEP and TFP confirmed that the relevant literature was included in the evidence synthesis. Unless they were identified by the literature search and met the article inclusion criteria (see Supplementary Appendix 3, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658), we did not review any unpublished data from product manufacturers, investigators, or the Food and Drug Administration (FDA) Adverse Event Reporting System.

We searched the literature for the 8 DMARDs and 9 biologic agents most commonly used for the treatment of RA. Literature was searched for 8 DMARDs: azathioprine, cyclosporine, hydroxychloroquine, leflunomide, methotrexate, minocycline, organic gold compounds, and sulfasalazine. Similar to 2008, azathioprine, cyclosporine, and gold were not included in the recommendations based on their infrequent use and lack of new data (Table 1). Literature was searched for 9 biologic agents: abatacept, adalimumab, anakinra, certolizumab pegol, etanercept, golimumab, infliximab, rituximab, and tocilizumab. Anakinra was not included in the recommendations due to infrequent use and lack of new data. Details of the bibliographic search strategy are listed in Supplementary Appendix 1 (available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658).

Literature search criteria and article selection.

Inclusion and exclusion criteria for review of article abstracts and titles. With the exception of assessment of TB, hepatitis, and vaccination (see below), studies were included if they met all of the following criteria: 1) original study in English language with an abstract, 2) observational studies (case–control or cohort) or intervention studies, 3) related to the treatment of RA with DMARDs or biologic agents, and 4) study duration of at least 6 months (see Supplementary Appendix 2, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658).

Studies were excluded if they met any of the following criteria: 1) the report was a meeting abstract, review article, or meta-analysis; 2) the study duration was less than 6 months; and 3) DMARDs or biologic agents were used for non-RA conditions (e.g., psoriatic arthritis, systemic lupus erythematosus) or non–FDA-approved use in health conditions other than RA (e.g., biologic agents in vasculitis) (see Supplementary Appendix 2, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658).

Selection criteria for articles reviewing efficacy/adverse events. Two reviewers independently screened the titles and abstracts of the 2,497 potential articles from the PubMed and Cochrane Library searches by applying the above selection method. Any disagreements were resolved by consultation with the lead reviewer (JAS). The lead author also reviewed all titles and abstracts to identify any that might have been overlooked. We identified 149 original articles from the 3 searches for full-text review. After excluding duplicates, 128 unique original articles were identified and the data were abstracted. This included 16 articles focused on DMARDs and 112 on biologic agents (98 on the 6 biologic agents assessed in the 2008 RA recommendations and 14 on certolizumab pegol, golimumab, and tocilizumab, 3 newer biologic agents that had been added since the 2008 recommendations) (see Supplementary Appendix 3, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658). A list of all included articles is provided in Supplementary Appendix 4 (available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658).

Additional literature searches for articles reviewing TB screening, hepatitis, and vaccination. Qualitative reviews of the literature were performed for these 3 topics (completed September 22, 2010). Similar to the strategy for the 2008 recommendations, literature searches were broadened to include case reports and case series of any size, review articles, and meta-analyses, plus inclusion of diseases other than RA. In addition, we included searches on the Centers for Disease Control and Prevention (CDC) web site (www.cdc.gov) for past and current recommendations regarding TB screening and vaccination in immunocompromised patients.

Agreement between reviewers for selection of studies for full-text retrieval. The kappa coefficients (agreement beyond chance) for independent selection of articles for full-text review by the 2 reviewers met or exceeded 0.60 (good) for DMARDs, 0.65 (very good) for the 6 biologic agents included in the 2008 ACR recommendations, and 0.84 (excellent) for a combination of certolizumab pegol, golimumab, and tocilizumab (9).

Full-text article review, data abstraction, data entry, and evidence report generation. The full text of each article was reviewed; data abstraction and entry were performed by reviewers using a standardized Microsoft Access database that was developed and used for data abstraction for the 2008 ACR RA recommendations. Two reviewers were assigned to abstract data on DMARDs (SB, DEF), rituximab (HA, ERV), and the rest of the biologic agents (AB, AJ). To ensure that the error rates were low and abstractions were similar, 26 articles related to biologic agents were dually abstracted by 2 abstractors (AB, AJ). The data entry errors were less than 3%. Entered data were further checked against raw data on biologic agents from the Cochrane systematic reviews (5–8). Following this comprehensive literature review, we developed an evidence report using the data abstracted from the published studies.

Development of clinical scenarios. Clinical scenarios were drafted by the investigators and the CEP, based on the updated evidence report. We used the same key determinant clinical thresholds and treatment decision branch points that were developed for the 2008 ACR RA treatment recommendations (1). Clinical scenarios were constructed based on permutations in the particular therapeutic considerations that reflected: 1) disease duration (early versus established RA), 2) disease activity (low, moderate, or high) (Tables 2 and 3 and Supplementary Appendix 5, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658),
Table 2. Definitions of key terms and key assumptions for clinical scenarios for the 2012 ACR recommendations update for the treatment of RA

<table>
<thead>
<tr>
<th>Definition</th>
<th>Key Terms</th>
</tr>
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<tbody>
<tr>
<td><strong>DMARDs†</strong></td>
<td>Hydroxychloroquine, leflunomide, methotrexate, minocycline, or sulfasalazine</td>
</tr>
<tr>
<td><strong>Non-methotrexate DMARDs</strong></td>
<td>Hydroxychloroquine, leflunomide, minocycline, or sulfasalazine</td>
</tr>
<tr>
<td><strong>DMARD combination therapy</strong></td>
<td>Combinations including 2 drugs, most of which are methotrexate based, with only a few exceptions (e.g., methotrexate + hydroxychloroquine, methotrexate + leflunomide, methotrexate + sulfasalazine, sulfasalazine + hydroxychloroquine), and triple therapy (methotrexate + hydroxychloroquine + sulfasalazine)</td>
</tr>
<tr>
<td><strong>Anti-TNF biologics</strong></td>
<td>Adalimumab, certolizumab pegol, etanercept, infliximab, or golimumab</td>
</tr>
<tr>
<td><strong>Non-TNF biologics</strong></td>
<td>Abatacept, rituximab, or tocilizumab</td>
</tr>
<tr>
<td><strong>Biologic agents</strong></td>
<td>Anti-TNF biologic or non-TNF biologic (8 biologic agents, excluding anakinra)</td>
</tr>
<tr>
<td><strong>Early RA</strong></td>
<td>RA disease duration &lt;6 months</td>
</tr>
<tr>
<td><strong>Established RA</strong></td>
<td>RA disease duration ≥6 months or meeting the 1987 ACR classification criteria (24)‡</td>
</tr>
<tr>
<td><strong>Disease activity</strong></td>
<td>Categorized as low, moderate, and high as per validated common scales (Table 3 and Supplementary Appendix 4, available in the online version of this article at <a href="http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658">http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658</a>) or the treating clinician’s formal assessment (26–32)</td>
</tr>
<tr>
<td><strong>Poor prognosis</strong></td>
<td>Presence of 1 or more of the following features: functional limitation (e.g., HAQ DI or similar valid tools), extraarticular disease (e.g., presence of rheumatoid nodules, RA vasculitis, Felty’s syndrome), positive rheumatoid factor or anti–cyclic citrullinated peptide antibodies (33–37), or bony erosions by radiograph (38–40)</td>
</tr>
<tr>
<td><strong>RA remission</strong></td>
<td>A joint ACR/EULAR task force defined remission as a tender joint count, swollen joint count, C-reactive protein (mg/dl) level, and patient global assessment of ≤1 each or a simplified Disease Activity Score of ≤3.3 (41)</td>
</tr>
<tr>
<td><strong>Child-Pugh classification</strong></td>
<td>Scoring system based upon the levels of albumin, total bilirubin, and prothrombin time, and the presence of ascites and encephalopathy. Patients with a score of 10 or more (in the class C category) have a prognosis with a 1-year survival rate of ~50%. Patients with class A or B have a better 5-year prognosis, with a survival rate of 70–80% (12)</td>
</tr>
<tr>
<td><strong>NYHA class III and IV</strong></td>
<td>NYHA class III includes patients with cardiac disease resulting in marked limitation of physical activity with less than ordinary physical activity causing fatigue, palpitation, dyspnea, or anginal pain, but no symptoms at rest. NYHA class IV includes patients with cardiac disease resulting in inability to carry on any physical activity without discomfort and symptoms of cardiac insufficiency or that the anginal syndrome may be present even at rest, which increases if any physical activity is undertaken (13)</td>
</tr>
<tr>
<td><strong>CDC-defined risk factors for latent TB infection</strong></td>
<td>Close contacts of persons known or suspected to have active TB; foreign-born persons from areas with a high incidence of active TB (e.g., Africa, Asia, Eastern Europe, Latin America, and Russia); persons who visit areas with a high prevalence of active TB, especially if visits are frequent or prolonged; residents and employees of congregate settings whose clients are at an increased risk for active TB (e.g., correctional facilities, long-term care facilities, and homeless shelters); health care workers who serve clients who are at an increased risk for active TB; populations defined locally as having an increased incidence of latent Mycobacterium tuberculosis infection or active TB, possibly including medically underserved, low-income populations, or persons who abuse drugs or alcohol; and infants, children, and adolescents exposed to adults who are at an increased risk for latent M tuberculosis infection or active TB (14)</td>
</tr>
</tbody>
</table>

**KEY ASSUMPTIONS**

1. Focus on common patients, not exceptional cases
2. Cost not considered; please see 2008 recommendations (1)
3. Alternate therapeutic choices taken into account
4. When a particular drug is not recommended, it does not imply that it is contraindicated
5. Examples of scenarios for which agreement is not achieved noted explicitly (see Supplementary Appendix 6, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658)§
6. Optimal dose of medication (as defined by the treating clinician) given for 3 months (in lieu of total duration of therapy) before therapy escalation or switching
7. Disease activity and prognosis assessments performed between 3 and 6 months after initiation or change in therapy, although can be assessed as early as 3 months
8. Assume that a clinical indication exists (based upon disease activity) for use of each treatment option

* ACR = American College of Rheumatology; RA = rheumatoid arthritis; DMARDs = disease-modifying antirheumatic drugs; anti-TNF = anti–tumor necrosis factor; HAQ = Health Assessment Questionnaire; DI = disability index; EULAR = European League Against Rheumatism; NYHA = New York Heart Association; CDC = Centers for Disease Control and Prevention; TB = tuberculosis.
† Azathioprine, cyclosporine, and gold were considered but not included due to their infrequent use in RA and/or the lack of new data since 2008.
‡ New classification criteria for RA (ACR/EULAR collaborative initiative) have been published (23); however, the evidence available for these new recommendations relied on the use of the 1987 ACR RA classification criteria, since literature review preceded the publication of the new criteria. We anticipate that in the near future, data using the new classification criteria may be available for evidence synthesis and formulating recommendations.
§ Agreement as defined by the RAND/UCLA Appropriateness Method.
3) current medication regimen, and 4) presence of poor prognostic factors (yes or no, as defined in the 2008 ACR recommendations). An example of a clinical scenario is: “The patient has active established RA and has failed an adequate trial of an Anti-TNF [anti–tumor necrosis factor] biologic because of adverse events. Is it appropriate to switch to another Anti-TNF biologic after failing etanercept?” (see Supplementary Appendix 6, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658). Scenarios included both new considerations and questions considered in the 2008 recommendations.

For this 2012 update, we used a modified Delphi process and obtained consensus (defined as ≥70% agreement) from the CEP for inclusion of relevant clinical scenarios based on 1) review of each of the previous 2008 scenarios and 2) review of newly developed scenarios to address switching between therapies. We provided CEP members with manuscript abstracts and requested full-text articles to help inform decisions.

The CEP members also recommended the following: 1) use of the FDA definitions of “serious” and “non-serious” adverse events (10), 2) exclusion of 3 DMARDs used very infrequently (i.e., cyclosporine, azathioprine, and gold; see above) or without additional relevant new data, and 3) exclusion of 1 biologic agent without additional relevant new evidence and with infrequent use (anakinra).

### Rating the appropriateness of clinical scenarios by the TFP

The TFP is referred to as the “panel” in the Methods and the recommendations that follow. For the first round of ratings we contacted panel members by e-mail and provided them with the evidence report, clinical scenarios, and rating instructions. We asked them to use the evidence report and their clinical judgment to rate the “appropriateness” of the clinical scenarios under consideration. The panelists individually rated each scenario permutation using a 9-point Likert appropriateness scale. A median score of 1 to 3 indicated “not appropriate” and 7 to 9 indicated “appropriate” for taking action defined in the scenario (2–4). For all eventual recommendations, the RAND/UCLA appropriateness panel score required a median rating of 7 to 9. Those scenario permutations with median ratings in the 4 to 6 range and those with disagreement among the panelists (i.e., one-third or more TFP members rating the scenario in the 1 to 3 range and one-third or more rating it in the 7 to 9 range) were classified as “uncertain.” At a face-to-face meeting with both the TFP and the CEP members on November 15, 2010, the anonymous first round of ratings by the panel, including dispersion of the scores, ranges, and median scores, was provided to the task force panelists.

The task force panelists agreed upon certain assumptions and qualifying statements on which they based their discussion and subsequent ratings of the scenarios (Table 2). A second round of ratings by panel members occurred after extensive in-person discussion of the prior ratings and review of the evidence supporting each scenario.

### Conversion of clinical scenarios to ACR RA treatment recommendations

After the TFP meeting was complete, recommendations were derived from directly transcribing the final clinical scenario ratings. Based on the ratings, scenario permutations were collapsed to yield the most parsimonious recommendations. For example, when ratings favored a drug indication for both moderate and high disease activity, one recommendation was given, specifying “moderate or high disease activity.” In most circumstances, the recommendations included only positive and not negative statements. For example, the recommenda-
tions focused on when to initiate specific therapies rather than when an alternate therapy should not be used. Most of the recommendations were formulated by drug category (DMARD, anti-TNF biologic, non-TNF biologic listed alphabetically within category), since in many instances, the ratings were similar for medications within a drug category. We specifically note instances where a particular medication was recommended but others in its group were not endorsed. Two additional community-based rheumatologists independently reviewed the manuscript and provided comments. CEP and TFP members reviewed and approved all final recommendations.

For each final recommendation, the strength of evidence was assigned using the methods from the American College of Cardiology (11). Three levels of evidence were specified: 1) level of evidence A: data were derived from multiple RCTs; 2) level of evidence B: data were derived from a single randomized trial or nonrandomized studies; and 3) level of evidence C: data were derived from consensus opinion of experts, case studies, or standards of care. The evidence was rated by 4 panel experts (JO and JMK, AFK and LWM, where each rated half of the evidence), and discrepancies were resolved by consensus.

Level C evidence often denoted a circumstance where medical literature addressed the general topic under discussion but it did not address the specific clinical situations or scenarios reviewed by the panel. Since many recommendations had multiple components (in most cases, multiple medication options), a range is sometimes provided for the level of evidence; for others, the level of evidence is provided following each recommendation.

ACR peer review of recommendations. Following construction of the recommendations, the manuscript was reviewed through the regular journal review process and by more than 30 ACR members serving on the ACR Guidelines Subcommittee, ACR Quality of Care Committee, and ACR Board of Directors.

Recommendations for the Use of DMARDs and Biologic Agents in Patients Who Qualify for Treatment of RA

This 2012 ACR recommendations update incorporates the evidence from systematic literature review synthesis and recommendations from 2008 (1) and rates updated and new clinical scenarios regarding the use of DMARDs and biologic agents for the treatment of RA. Terms used in the recommendations are defined in Table 2. The 2012 recommendations are listed in the 4 sections below and in the following order:

1. Indications for and switching DMARDs and biologic agents: early RA (indications, Figure 1) followed by established RA (indications and switching, Figure 2), along with details of the level of evidence supporting these recommendations (see Supplementary Appendix 7, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN) 2151-4658)

2. Use of biologic agents in patients with hepatitis, malignancy, or CHF who qualify for RA management (Table 4)

3. Screening for TB in patients starting or currently receiving biologic agents as part of their RA therapy (Figure 3)

4. Vaccination in patients starting or currently receiving DMARDs or biologic agents as part of their RA therapy (Table 5)

In the figures, decision points are shown as diamonds and actions to be taken by the health care provider are shown as rectangles. The recommendations in the text below and in Tables 4 and 5 represent the results of the 2012 update only, whereas Figures 1–3 also incorporate some of the 2008 ACR RA recommendations that did not change (1). Areas of uncertainty by the panel (that did not lead to recommendations) are noted in Supplementary Appendix 8 (available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN) 2151-4658).

1. Indications for starting, resuming, adding, or switching DMARDs or biologic agents. We first describe a recommendation targeting remission or low disease activity in RA (section 1A). This is followed by recommendations for DMARD or biologic agent use in early RA (section 1B). Next, we provide recommendations for initiating and switching between DMARDs and biologic agents in established RA (section 1C).

1A. Target low disease activity or remission. The panel recommends targeting either low disease activity (Table 3) or remission (Table 2) in all patients with early RA (Figure 1; level of evidence C) and established RA (Figure 2; level of evidence C) receiving any DMARD or biologic agent.

1B. Early RA (disease duration <6 months). In patients with early RA, the panel recommends the use of DMARD monotherapy both for low disease activity and for moderate or high disease activity with the absence of poor prognostic features (Figure 1; level of evidence A–C) (details are shown in Supplementary Appendix 7, available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658).

In patients with early RA, the panel recommends the use of DMARD combination therapy (including double and triple therapy) in patients with moderate or high disease activity plus poor prognostic features (Figure 1; level of evidence A–C).

In patients with early RA, the panel also recommends the use of an anti-TNF biologic with or without methotrexate in patients who have high disease activity with poor prognostic features (Figure 1; level of evidence A and B). Infliximab is the only exception and the recommendation is to use it in combination with methotrexate, but not as monotherapy.

1C. Established RA (disease duration ≥6 months or meeting the 1987 ACR RA classification criteria). The remainder of panel recommendations regarding indications for DMARDs and biologic agents are for patients with established RA. The 3 subsections below define recommendations for initiating and switching therapies in established RA (Figure 2). Where the prognosis is not mentioned, the recommendation to use/switch to a DMARD or a biologic agent applies to all patients, regardless of prognostic features.
Initiating and switching among DMARDs.

If after 3 months of DMARD monotherapy (in patients without poor prognostic features), a patient deteriorates from low to moderate/high disease activity, then methotrexate, hydroxychloroquine, or leflunomide should be added (rectangle A of Figure 2; level of evidence A and B).

If after 3 months of methotrexate or methotrexate/DMARD combination, a patient still has moderate or high disease activity, then add another non-methotrexate DMARD or switch to a different non-methotrexate DMARD (rectangle B of Figure 2; level of evidence B and C).

Switching among biologic agents due to lack of benefit or loss of benefit.

If a patient still has moderate or high disease activity after 3 months of anti-TNF biologic therapy and this is due to a lack or loss of benefit, switching to another anti-TNF biologic or a non-TNF biologic is recommended (rectangles F and G of Figure 2; level of evidence C).

If a patient still has moderate or high disease activity after 6 months of a non-TNF biologic and the failure is due to a lack or loss of benefit, switch to another non-TNF biologic or an anti-TNF biologic (rectangles F and G of Figure 2; level of evidence B and C). An assessment period of 6 months was chosen rather than 3 months, due to the anticipation that a longer time may be required for efficacy of a non-TNF biologic.

Switching among biologic agents due to harms/adverse events.

If a patient has high disease activity after failing an anti-TNF biologic because of a serious adverse event, switch to a non-TNF biologic (rectangle E of Figure 2; level of evidence C).

If a patient has moderate or high disease activity after failing an anti-TNF biologic because of a nonserious ad-
Figure 2. 2012 American College of Rheumatology (ACR) recommendations update for the treatment of established rheumatoid arthritis (RA), defined as a disease duration ≥6 months or meeting the 1987 ACR classification criteria. Depending on a patient’s current medication regimen, the management algorithm may begin at an appropriate rectangle in the figure, rather than only at the top of the figure. Disease-modifying antirheumatic drugs (DMARDs) include hydroxychloroquine (HCQ), leflunomide (LEF), methotrexate (MTX), minocycline, and sulfasalazine (therapies are listed alphabetically; azathioprine and cyclosporine were considered but not included). DMARD monotherapy refers to treatment in most instances with HCQ, LEF, MTX, or sulfasalazine; in few instances, where appropriate, minocycline may also be used. Anti–tumor necrosis factor (anti-TNF) biologics include adalimumab, certolizumab pegol, etanercept, infliximab, and golimumab. Non-TNF biologics include abatacept, rituximab, or tocilizumab (therapies are listed alphabetically). For the level of evidence supporting each recommendation, please see Supplementary Appendix 7 (available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658).

* Definitions of disease activity are discussed in Tables 2 and 3 and Supplementary Appendix 4 (available in the online version of this article at http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)2151-4658) and were categorized as low, moderate, or high.

† Features of poor prognosis included the presence of 1 or more of the following: functional limitation (e.g., Health Assessment Questionnaire score or similar valid tools), extraarticular disease (e.g., presence of rheumatoid nodules, RA vasculitis, Felty’s syndrome), positive rheumatoid factor or anti–cyclic citrullinated peptide antibodies (33–37), and bony erosions by radiograph (38).

‡ Combination DMARD therapy with 2 DMARDs, which is most commonly MTX based, with few exceptions (e.g., MTX + HCQ, MTX + LEF, MTX + sulfasalazine, sulfasalazine + HCQ), and triple therapy (MTX + HCQ + sulfasalazine).

§ Reassess after 3 months and proceed with escalating therapy if moderate or high disease activity in all instances except after treatment with a non-TNF biologic (rectangle D), where reassessment is recommended at 6 months due to a longer anticipated time for peak effect.

¶ LEF can be added in patients with low disease activity after 3–6 months of minocycline, HCQ, MTX, or sulfasalazine.

# If after 3 months of intensified DMARD combination therapy or after a second DMARD has failed, the option is to add or switch to an anti-TNF biologic.

** Serious adverse events were defined per the US Food and Drug Administration (FDA; see below); all other adverse events were considered nonserious adverse events.

†† Reassessment after treatment with a non-TNF biologic is recommended at 6 months due to anticipation that a longer time to peak effect is needed for non-TNF compared to anti-TNF biologics.

‡‡ Any adverse event was defined as per the US FDA as any undesirable experience associated with the use of a medical product in a patient. The FDA definition of serious adverse event includes death, life-threatening event, initial or prolonged hospitalization, disability, congenital anomaly, or an adverse event requiring intervention to prevent permanent impairment or damage.
Table 4. 2012 American College of Rheumatology recommendations update for the use of biologic agents in patients otherwise qualifying for the rheumatoid arthritis treatment with a history of hepatitis, malignancy, or congestive heart failure*  

<table>
<thead>
<tr>
<th>Comorbidity/clinical circumstance</th>
<th>Recommended</th>
<th>Not recommended</th>
<th>Level of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hepatitis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepatitis C</td>
<td>Etanercept</td>
<td>Any biologic agent</td>
<td>C</td>
</tr>
<tr>
<td>Untreated chronic hepatitis B or with treated chronic hepatitis B with Child-Pugh class B and higher†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Malignancy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Treated solid malignancy &gt;5 years ago or treated nonmelanoma skin cancer &gt;5 years ago</td>
<td>Any biologic agent</td>
<td></td>
<td>C</td>
</tr>
<tr>
<td>Treated solid malignancy within the last 5 years or treated nonmelanoma skin cancer within the last 5 years‡</td>
<td>Rituximab</td>
<td></td>
<td>C</td>
</tr>
<tr>
<td>Treated skin melanoma‡</td>
<td>Rituximab</td>
<td></td>
<td>C</td>
</tr>
<tr>
<td>Treated lymphoproliferative malignancy</td>
<td>Rituximab</td>
<td></td>
<td>C</td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NYHA class III/IV and with an ejection fraction of ≤50%§</td>
<td>Anti-TNF biologic</td>
<td></td>
<td>C</td>
</tr>
</tbody>
</table>

* For definitions and key terms, please refer to Table 2. NYHA = New York Heart Association; anti-TNF = anti-tumor necrosis factor.  
† Therapy defined as an antiviral regimen deemed appropriate by an expert in liver diseases. The Child-Pugh classification liver disease scoring system is based on the presence of albumin, ascites, total bilirubin, prothrombin time, and encephalopathy. Patients with a score of 10 or more (in the class C category) have a prognosis with 1-year survival being ~50%. Patients with class A or B have a better prognosis of 5 years, with a survival rate of 70–80% (12).  
‡ Little is known about the effects of biologic therapy on solid cancers treated within the past 5 years, due to exclusion of these patients from most randomized controlled trials.  
§ NYHA class III = patients with cardiac disease resulting in marked limitation of physical activity. They are comfortable at rest. Less than ordinary physical activity causes fatigue, palpitation, dyspnea, or anginal pain. NYHA class IV = patients with cardiac disease resulting in inability to carry out any physical activity without discomfort. Symptoms of cardiac insufficiency or of anginal syndrome may be present even at rest. If any physical activity is undertaken, discomfort is increased (13).  

verse event, switch to another anti-TNF biologic or a non-TNF biologic (rectangle F of Figure 2; level of evidence B and C).

If a patient has moderate or high disease activity after failing a non-TNF biologic because of an adverse event (serious or nonserious), switch to another non-TNF biologic or an anti-TNF biologic (rectangle F of Figure 2; level of evidence C).

2. Use of biologic agents in RA patients with hepatitis, malignancy, or CHF, qualifying for more aggressive treatment (level of evidence C for all recommendations).

Hepatitis B or C. The panel recommends that etanercept could potentially be used in RA patients with hepatitis C requiring RA treatment (Table 4). The panel also recommends not using biologic agents in RA patients with untreated chronic hepatitis B (disease not treated due to contraindications to treatment or intolerable adverse events) and in RA patients with treated chronic hepatitis B with Child-Pugh class B and higher (Table 4; for details of Child-Pugh classification, see Table 2) (12). The panel did not make recommendations regarding the use of any biologic agent for treatment in RA patients with a history of hepatitis B and a positive hepatitis B core antibody.

Malignancies. For patients who have been treated for solid malignancies more than 5 years ago or who have been treated for nonmelanoma skin cancer more than 5 years ago, the panel recommends starting or resuming any biologic agent if those patients would otherwise qualify for this RA management strategy (Table 4).

The panel only recommends starting or resuming rituximab in RA patients with: 1) a previously treated solid malignancy within the last 5 years, 2) a previously treated nonmelanoma skin cancer within the last 5 years, 3) a previously treated melanoma skin cancer, or 4) a previously treated lymphoproliferative malignancy. Little is known about the effects of biologic therapy in patients with a history of a solid cancer within the past 5 years owing to the exclusion of such patients from participation in clinical trials and the lack of studies examining the risk of recurrent cancer in this subgroup of patients.

CHF. The panel recommends not using an anti-TNF biologic in RA patients with CHF that is New York Heart Association (NYHA) class III or IV and who have an ejection fraction of 50% or less (Table 4) (13).

3. TB screening for biologic agents (level of evidence C for all recommendations except for initiation of biologic agents in patients being treated for latent TB infection [LTBI], where the level of evidence is B). The panel recommends screening to identify LTBI in all RA patients being considered for therapy with biologic agents, regardless of the presence of risk factors for LTBI (diamond A of Figure 3) (14). It recommends that clinicians assess the patient’s medical history to identify risk factors for TB (specified by the CDC) (Table 2).

The panel recommends the tuberculin skin test (TST) or interferon-γ-release assays (IGRAs) as the initial test in all RA patients starting biologic agents, regardless of risk factors for LTBI (diamond A of Figure 3). It recommends the use of the IGRA over the TST in patients who had previ-
Figure 3. 2012 American College of Rheumatology recommendations update for tuberculosis (TB) screening with biologic agent use. Depending on a patient’s current therapy, the management may begin at an appropriate rectangle in the figure, rather than only at the top of the figure. The level of evidence supporting each recommendation for TB reactivation was “C,” except for initiation of biologic agents in patients being treated for latent TB infection, where the level of evidence was “B.”

* Anergy panel testing is not recommended.
† Interferon-γ-release assay (IGRA) is preferred if the patient has a history of BCG vaccination.
‡ Risk factors for TB exposure are defined based on a publication from the US Centers for Disease Control and Prevention as: close contacts of persons known or suspected to have active TB; foreign-born persons from areas that have a high incidence of active TB (e.g., Africa, Asia, Eastern Europe, Latin America, and Russia); persons who visit areas with a high prevalence of active TB, especially if visits are frequent or prolonged; residents and employees of congregate settings whose clients are at an increased risk for active TB (e.g., correctional facilities, long-term care facilities, and homeless shelters); health care workers who serve clients who are at an increased risk for active TB; populations defined locally as having an increased incidence of latent Mycobacterium tuberculosis infection or active TB, possibly including medically underserved, low-income populations, or persons who abuse drugs or alcohol; and infants, children, and adolescents exposed to adults who are at an increased risk for latent M tuberculosis infection or active TB (14).
§ If the patient is immunosuppressed and false-negative results are more likely, consider repeating screening test(s) with tuberculin skin test (TST) or IGRA.
¶ Chest radiograph may also be considered when clinically indicated in patients with risk factors, even with a negative repeat TST or IGRA.
# Obtain respiratory (e.g., sputum, bronchoalveolar lavage fluid) or other samples as clinically appropriate for acid-fast bacilli (AFB) smear and culture and consider referral to a TB specialist for further evaluation and treatment.
** In a patient diagnosed with latent or active TB, consider referral to a specialist for the recommended treatment.
†† Patients who test positive for TST or IGRA at baseline often remain positive for these tests even after successful treatment of TB. These patients need monitoring for clinical signs and symptoms of recurrent TB disease, since repeating tests will not allow help in diagnosis of recurrent TB.
viously received a BCG vaccination, due to the high false-positive test rates for TST (Figure 3).

The panel recommends that RA patients with a positive initial or repeat TST or IGRA should have a chest radiograph and, if suggestive of active TB, a subsequent sputum examination to check for the presence of active TB (diamonds B and C of Figure 3). RA patients with a negative screening TST or IGRA may not need further evaluation in the absence of risk factors and/or clinical suspicion for TB. Since patients with RA may have false-negative TST or IGRA results due to immunosuppression, a negative TST or IGRA should not be interpreted as excluding the possibility that a patient has LTBI. Accordingly, in immunosuppressed RA patients with risk factors for LTBI and negative initial screening tests, the panel recommends that a repeat TST or IGRA could be considered 1–3 weeks after the initial negative screening (diamond A of Figure 3).

If the RA patient has active or latent TB based on the test results, the panel recommends appropriate antitubercular treatment and consideration of referral to a specialist. Treatment with biologic agents can be initiated or resumed after 1 month of latent TB treatment with antitubercular medications and after completion of the treatment of active TB, as applicable (Figure 3).

The panel recommends annual testing in RA patients who live, travel, or work in situations where TB exposure is likely while they continue treatment with biologic agents (diamond D of Figure 3). Patients who test positive for TST or IGRA at baseline can remain positive for these tests even after successful treatment of TB. These patients need monitoring for clinical signs and symptoms of recurrent TB, since repeating tests will not help in the diagnosis of recurrent TB.

4. Vaccination in patients starting or currently receiving DMARDs or biologic agents as part of their RA therapy (level of evidence C for all recommendations). The panel recommends that all killed (pneumococcal, influenza intramuscular, and hepatitis B), recombinant (human papillomavirus [HPV] vaccine for cervical cancer), and live attenuated (herpes zoster) vaccinations should be undertaken before starting a DMARD or a biologic agent (Table 5).

It also recommends that, if not previously done, vaccination with indicated pneumococcal (killed), influenza intramuscular (killed), hepatitis B (killed), and HPV vaccine (recombinant) should be undertaken in RA patients already taking a DMARD or a biologic agent (Table 5).

The panel recommends vaccination with herpes zoster vaccine in RA patients already taking a DMARD. All vaccines should be given based on age and risk, and physicians should refer to vaccine instructions and CDC recommendations for details about dosing and timing issues related to vaccinations.

**Discussion**

We updated the 2008 ACR RA recommendations for the treatment of RA (1) using the scientific evidence and a rigorous evidence-based group consensus process. The 2012 update addresses the use of DMARDs and biologic agents, switching between therapies, the use of biologic agents in high-risk patients, TB screening with the use of biologic agents, and vaccination in patients with RA receiving DMARDs or biologic agents.

The rigorous process to which we referred above included a comprehensive updated literature review, data

<p>| Table 5. 2012 American College of Rheumatology recommendations update regarding the use of vaccines in patients with RA starting or currently receiving DMARDs or biologic agents* |</p>
<table>
<thead>
<tr>
<th>DMARD monotherapy</th>
<th>Combination DMARDs§</th>
<th>Anti-TNF biologics¶</th>
<th>Non-TNF biologics#</th>
<th>Recombinant vaccine</th>
<th>Killed vaccines</th>
<th>Live attenuated vaccine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pneumococcal†</td>
<td>Influenza (intramuscular)</td>
<td>Hepatitis B#</td>
<td>Human papillomavirus</td>
<td>Herpes zoster</td>
<td>Not recommended**</td>
<td>Not recommended**</td>
</tr>
<tr>
<td>Before initiating therapy</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>While already taking therapy</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>

* Evidence level was C for all of the vaccination recommendations. For definitions and key terms, please refer to Table 2. DMARDs = disease-modifying antirheumatic drugs; √ = recommend vaccination when indicated (based on age and risk); anti-TNF = anti-tumor necrosis factor.
† The Centers for Disease Control and Prevention also recommends a one-time pneumococcal revaccination after 5 years for persons with chronic conditions such as rheumatoid arthritis (RA). For persons ages ≥65 years, one-time revaccination is recommended if they were vaccinated ≥5 years previously and were age ≥65 years at the time of the primary vaccination.
‡ If hepatitis risk factors are present (e.g., intravenous drug abuse, multiple sex partners in the previous 6 months, health care personnel).
§ DMARDs include hydroxychloroquine, leflunomide, methotrexate, minocycline, and sulfasalazine (listed alphabetically) and combination DMARD therapy included double (most methotrexate based, with few exceptions) or triple therapy (hydroxychloroquine + methotrexate + sulfasalazine).
¶ Anti-TNF biologics include adalimumab, certolizumab pegol, etanercept, golimumab, and infliximab (listed alphabetically).
# Non-TNF biologics include abatacept, rituximab, and tocilizumab (listed alphabetically).
** According to the RAND/UCLA Appropriateness Method, panel members judged it as “not appropriate” and therefore it qualifies as “not recommended” (median score on appropriateness scale was 1).
review by a panel of international experts, and use of a well-accepted validated process for developing recommendations (2–4).

Because we used the same method for this update as the 2008 ACR RA recommendations, we were able to incorporate the evidence from the 2008 process and comprehensively update the recommendations. Consistent with the common need to extrapolate from clinical experience in the absence of higher-tier evidence, many of these new recommendations (approximately 79%) were associated with level C evidence.

These recommendations aim to address common questions facing both patients with RA and the treating health care providers. Since the recommendations were derived considering the “common patients, not exceptional cases,” they are likely to be applicable to a majority of RA patients, although not all patients. As always, clinical judgment must be used. The emergence of several new therapies for RA in the last decade has led to great excitement in the field of rheumatology as well as provided patients and health care providers with multiple options for treatment.

The 2008 recommendations and 2012 update attempt to simplify the treatment algorithms for patients and providers. These recommendations provide clinicians with choices for treatments of patients with active RA, both in early and established disease phases.

Recommendations also provide guidance regarding treatment choices in RA patients with comorbidities such as hepatitis, CHF, and malignancy. In particular, the risk for TB reactivation has become an increasingly common concern for clinicians and patients treating RA patients with biologic agents. The algorithm recommended provides a comprehensive approach for many RA patients. Due to an increasing awareness of risk of preventable diseases such as influenza and pneumonia (especially in the elderly), immunizations are very important in patients with RA. Several recommendations address this important aspect of vaccination of RA patients. Because these recommendations were heavily informed by CDC guidance and minimal additional information was found in the broader literature search, our TB screening and vaccination recommendations are concordant with the CDC recommendations.

The goal for each RA patient should be low disease activity or remission. In ideal circumstances, RA remission should be the target of therapy, but in others, low disease activity may be an acceptable target. But for other patients, the decision about what the target should be for each patient is appropriately left to the clinician caring for each RA patient, in the context of patient preferences, comorbidities, and other individual considerations. Therefore, this article does not recommend a specific target for all patients. Of note, the panel recommended a more aggressive treatment in patients with early RA than in the 2008 ACR recommendations. We speculate that this may be related to several reasons: 1) the expectation that the earlier the treatment the better the outcome, 2) the thought that joint damage is largely irreversible so prevention of damage is an important goal, and 3) the data that early intensive therapy may provide the best opportunity to preserve physical function and health-related quality of life and reduce work-related disability (15–22).

As with all recommendations, these recommendations apply to common clinical scenarios, and only a clinician’s assessment in collaboration with the patient allows for the best risk–benefit analysis on a case-by-case basis. These recommendations cannot adequately convey all uncertainties and nuances of patient care in the real world. For example, the panel did not vote on the possibility of temporarily holding biologic therapy to facilitate administration of the live herpes zoster vaccine among older patients and then resuming the biologic agent shortly thereafter. All recommendations were based on scientific evidence coupled with our formal group process rather than only the approved indications from regulatory agencies. Although new classification criteria for RA (ACR/European League Against Rheumatism collaborative initiative) were published in September 2010 (23), the studies evaluated for the 2012 recommendations relied on the use of the 1987 ACR RA classification criteria (24) because our literature review preceded the publication of the new criteria.

The need to create recommendations that cover a comprehensive array of relevant clinical decisions has led to many recommendations that combine literature-based data and expert opinion, and thus are labeled as level C evidence. For example, rituximab was recommended as appropriate in patients with previously treated solid malignancy within the last 5 years or a previously treated nonmelanoma skin cancer within the last 5 years, which is a level C recommendation since the evidence is based on clinical trial extensions and observational data. It is important to note that the limited evidence available supporting this recommendation comes primarily from non-RA populations that included cancer patients. In addition, the panel ratings did not achieve the level of appropriateness needed to recommend other biologic therapies in this circumstance since most of the panelists’ ratings were “uncertain.” Like many of the other recommendations put forth, this recommendation was grounded, in part, on expert consensus and serves to highlight an important evidence gap in RA management.

In some cases the panelists did not make a specific recommendation statement. This occurred when ratings reflected uncertainty over a particular potential clinical scenario or when there was inability to reach consensus. In these cases, given a lack of clear evidence or clear consensus, therapeutic decisions are best left to the careful consideration of risks/benefits by the individual patient and physician. These areas could be the subject of future research agendas and recommendations updates.

We anticipate that in the future, data using the new RA classification criteria (23) may be available for evidence synthesis and formulating recommendations. Recommendations regarding the use of other antiinflammatory medications, such as nonsteroidal antiinflammatory drugs and intraarticular and oral corticosteroids, and nonpharmacologic therapies (such as physical and occupational therapies) were not within the purview of this update, although these may be important components of RA treatment and could also be included in separate reviews or recommendations. For example, recommendations related to glucocorticoid use in RA have been published by other professional organizations (25). In the future, the ACR may decide to develop broader RA guidelines that include ther-
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