Economic evaluation of anti-TNF agents for patients with rheumatoid arthritis in Greece

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Objectives: We aimed to estimate the total mean annual treatment cost of different therapy options for patients with moderate-to-severe rheumatoid arthritis (RA) in Greece.

Methods: A cost-minimization approach was adopted. An economic model was developed to estimate the direct costs of the three widely used treatments within a 1-year time horizon, from a health care payer perspective, either for new or for existing patients. Data on resource use, dose escalation, and frequency of therapy were based on a nationwide field survey of rheumatologists. Other analyses were also undertaken based on evidence from the literature. Total cost comprised the cost of drugs, administration, and hospital day care visits. Unit cost data were obtained from the price bulletin and the government gazettes issued by the Ministry of Health. Due to the short time horizon of the study, the cost was not discounted.

Results: The mean annual total cost per new (or per existing) responder patient on etanercept was estimated at €9,845 (€9,840), and the total cost on etanercept/methotrexate (MTX) was estimated at €9,857 (€9,852). Therapy with etanercept had lower annual cost relative to adalimumab and infliximab. On an annual basis, it was estimated that the difference between etanercept monotherapy and adalimumab monotherapy was €544 (€1,323). Similarly, the difference between etanercept/MTX and infliximab/MTX was €1,871 (€1,490) and €543 (€1,323), respectively, relative to adalimumab/MTX. Results remained constant under other scenario analyses undertaken.

Conclusion: In the real-life practice setting in Greece, where dose intensity and frequency differences occur, etanercept alone or in combination with MTX, if prescribed as per label, represents the option with lower annual cost per patient when compared with adalimumab or infliximab in patients with RA. These results hold true as long as the assumptions and data used in the analysis remain stable and may alter if any of the underlying parameters, such as drug price, change.

Keywords: etanercept, dose escalation, economic analysis

Introduction

Rheumatoid arthritis (RA) is an inflammatory disease characterized by persistent synovitis, systemic inflammation, and autoantibodies. Several studies have reported overall RA prevalence rates of up to 0.67% of the total population, and they have also indicated that it is more common amongst women rather than men. Several cost-of-illness studies undertaken in a variety of settings and countries, including the USA, Canada, the Netherlands, Sweden, and Germany, have indicated that RA imposes a significant direct and indirect burden on patients, their families, and society overall. This has been also confirmed by a recent review of the literature that considered 127 relevant articles. Boonen and Severens combined results coming...
from 26 cost-of-illnesses studies and estimated that the total mean annual cost per RA patient was about €21,900, which is a noticeable figure.18

The European League Against Rheumatism (EULAR) and the American College of Rheumatology (ACR) have developed treatment guidelines for RA patients.19,20 These recommend the use of biological disease modifying antirheumatic drugs (DMARDS) after the failure of conventional DMARDS or as a first-line therapy option for patients with poor prognostic factors.19,20 Similar recommendations have been elaborated for the case of Greece.21,22 Existing evidence suggests that timely introduction of biologic DMARDS may improve clinical outcomes, induce remission, prevent radiographic progression, improve quality of life, and reduce lost work days or early retirement.23 The HOPE study reported that reduced or delayed access to appropriate RA treatment can cause considerable economic burden to patients and society, due to the significant costs associated with the higher consumption of health services needed for treating advanced-stage disease, higher productivity losses, and disability benefits.24

In addition to timely access to therapy, treatment costs are dependent on the intensity and frequency of drug doses prescribed in real-life settings. Furthermore, the mode of administration of the therapeutic options differs and hence, their resource utilization. Therefore, from a payer’s perspective, the actual cost of therapy depends on the type of medication prescribed and its unit price on the one hand, and on the other hand, the frequency and dose prescribed in real-life settings. For these reasons, several analyses reported in the literature have attempted to compare the alternative therapies in terms of their costs, based on data from clinical practice, and in fact, they have indicated profound differences amongst the alternative therapies.25–27 Given the aforementioned, to make rational comparisons and subsequent choices between alternative therapies for RA on the basis of economic evidence, the actual prescribed doses in real-life settings must be considered and therapies should be evaluated based on their actual, rather than theoretical cost.

Greece is going through a very deep and sustained economic crisis, which has eliminated a significant portion of its gross domestic product (GDP) and which has had profound social, financial, political, health care, and health implications.28 An unprecedented reform is currently being implemented in the health care sector in order to increase efficiency in the funding and provision of services. The health care reforms in the domain of pharmaceuticals are attempting to revise the pricing and reimbursement system, to introduce prescription utilization and dispensing controls, and to restrain overall pharmaceutical expenditure through fixed budgets, price-volume, risk-sharing agreements, and similar measures. RA therapies represent one of the leading classes of products in terms of cost. Within this setting, and taking into account the aforementioned timing and cost determinants of RA therapy, it is paramount for payers to ensure that RA patients receive optimum therapy in terms of effectiveness but also, in terms of efficiency and cost-effectiveness. Given the number of patients and the size of budget concerned, the aforementioned should be a very important payer policy objective.

EOPYY, the National Organization for Health Services Provision, is the single national payer that was created from the merging of the health branches of all sickness funds in the country, and it provides coverage to all patients in the country. For the EOPYY, it is important to know the health and economic outcomes associated with the use of different biologic treatments for RA and hence, the rational for the present study, which was designed to provide some insights into this area.

Methods
Etanercept, infliximab, and adalimumab, represent three widely used anti-TNF (Tumor Necrosis Factor), available for the management of RA patients.39 The aim of the present study was to estimate the annual mean cost of treating an RA patient with etanercept, infliximab, and adalimumab either as monotherapy, when appropriate, or in combination with methotrexate (MTX), in the Greek health care setting. Although there is no direct comparative evidence on the relative efficacy of the three agents, an indirect comparison of randomized placebo-controlled trials undertaken in the context of a systematic review and meta-analysis of the literature indicated that there may be no substantial differences amongst them in terms of efficacy and safety.40 Hence the therapies were considered equivalent in terms of health outcomes, and therefore a cost-minimization analysis was undertaken on this grounds. This is the appropriate economic evaluation method in such cases. In this context, only the therapy cost associated with the use of each of the selected alternative treatments for RA in Greece was quantified. A Microsoft Excel® economic model was developed to estimate the cost associated with actual prescribed drug and administration for each alternative therapy. The time horizon of the model was limited to 1 year. The cost of adverse events associated with the different comparators was not considered, as no significant differences amongst comparators were found.40
The model simulated patients during the course of a year and quantified the cost of resource consumption. In each therapy option, the model simulated results for two different groups of patients starting therapy, “responders” and “nonresponders”. Responders were identified as those who attained the minimum improvement criterion for RA, ACR 20, defined by the ACR as a 20% improvement in the counts of the number of tender and swollen joints and at least three items from the following: observer evaluation of overall disease activity; patient evaluation of overall disease activity; patient evaluation of pain; a score of physical disability; and improvements in blood acute phase responses.

The model assumed that treated patients who had attained an ACR 20, continued to maintain this improvement while on therapy with a specific biologic agent. The model assumed that nonresponders were evaluated and dropped out from therapy due to lack of efficacy after a period of 12 weeks. Results were estimated for patients starting therapy for the first time and for existing patients who continued on therapy for another year. The prices of the drugs used in the model were based on the price bulletin issued by the Ministry of Health. The cost of drug therapy was calculated based on the available package size and cost and the required dose per patient. The present analysis, as mentioned above, was conducted from a payer’s perspective. This analysis includes the cost of pharmaceuticals, expressed in hospital price, and the cost of administration related to day care hospitalization (ie, infusion for intravenous therapies) or physician visit (ie, for outpatient subcutaneous therapies).

The main input parameters driving the results of the model were the dose and frequency of prescribed drugs. The MTX dose was assumed at 15 mg per os per week. Anti-TNF recommended doses are: etanercept 50 mg, administered through a subcutaneous injection once a week, or 25 mg, administered by subcutaneous injections twice per week; infliximab 3 mg/kg at week 0, 2, and 6, and every 8 weeks thereafter, administered by intravenous injection; and adalimumab 40 mg every other week, administered by subcutaneous injection. A possibility for dose escalation and/or intensification is possible in the case of infliximab and adalimumab, as referenced in their respective Summary of Product Characteristics (SPC). Hence, after initial therapy, a possibility for dose and frequency increase was incorporated into the model, for the latter two therapy options.

Data concerning the dose escalation were obtained from a nationwide physician survey, performed during a 2-month period between July and August of 2013. The data were collected through 10-minute telephone interviews with rheumatologists. Each participating physician was interviewed based on a structured questionnaire designed to collect information regarding the demographics of managed patients and the followed treatment patterns. Simple random sampling was used to select the interviewees. Specifically, participants were randomly selected through a publicly available database, which includes contact details for all registered members of the Hellenic Society of Rheumatology (http://www.ere.gr). Sample size calculations were done prior to the study to make sure that the interviewed participants would constitute a representative sample of the rheumatologist population in Greece. A screening question was used to identify physicians who had used biologic agents for the treatment of RA in the last 12 months prior to interview. A weight was attached to each responder, based on the patient group size. These weights were then used to aggregate the responses of the interviewed sample.

The unit cost per item used in the model is shown in Table 1. Due to time horizon of the analysis, discounting was unnecessary. In studies such as the present one, there are a few uncertainties present either because of the nature of the assumptions necessary to undertake the analysis or because of the uncertainly and variability of the data utilized. Based on the data collected, an empirical distribution was estimated, and it was then used to run a probabilistic sensitivity analysis. Specifically, the dose and frequency of escalation variables were associated with the empirical distributions; then, in each of 10,000 simulations, a different variable was drawn at random from the distribution, and the results were recalculated. The set of obtained results was then used to estimate uncertainty intervals. Moreover, to further test the robustness of the results, alternative scenarios were tested, where data on dose escalation were based on evidence derived from the literature. In this context, the economic analysis was undertaken two more times. In the first scenario, the escalation rate was: 2.5% for etanercept, 9.6% for adalimumab, and 35% for infliximab according to evidence from the DART study, and in the second scenario, the escalation rate was

<table>
<thead>
<tr>
<th>Table 1 Unit cost per item used in the model</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Description</strong></td>
</tr>
<tr>
<td>Etanercept 50 mg/1 mL ×4 prefilled syringe</td>
</tr>
<tr>
<td>Infliximab 100 mg/vial</td>
</tr>
<tr>
<td>Adalimumab 40 mg</td>
</tr>
<tr>
<td>Methotrexate 2.5 mg/tablet, 100 tablet bottle</td>
</tr>
<tr>
<td>Physician office visit*</td>
</tr>
<tr>
<td>Day care hospitalization*</td>
</tr>
</tbody>
</table>

0%, 0%, and 55% respectively, according to evidence from the Hellenic Registry of Biologics."\(^{46}\)

**Results**

The sample of participating physicians in the survey consisted of 96 rheumatologists who were based in Athens (69%), Thessaloniki (13%), and other urban areas (18%), such as Patras, Larissa, and Heraklion. Assuming from the published literature a prevalence of disease of 0.67\(^{\%}\)^\(^{5}\), the moderate/severe RA patient population in Greece may be estimated at around 27,000 cases. Altogether, the physicians included in the study reported treating 4,015 patients in the year of the study, and thus it may be inferred that they were treating about 15\% of all moderate/severe patients in Greece, which is a significant sample size. About one-third (33\%) of the patients managed by those participating physicians were male and the remaining (67\%) were female.

For patient cases on infliximab who experienced lack of response, either the dose was increased from 3 mg/kg to 4.5 mg/kg or the interval reduced from 8 to 6 weeks, and these figures were used in the baseline analysis.\(^{37}\) For patient cases on adalimumab who experience lack of response, the dose increase was from 40 mg to 73.9 mg and the dose interval was reduced from every other week to every week, and this is assumed in the analysis.\(^{26}\) Moreover, as seen in Table 2, the initial prescribed dose of infliximab in about one-third of patients was not the one recommended in the SPC but rather, was higher in dose intensity (14.3\%) or frequency (11.9\%), or in a combination of the two (3.6\%). As seen in Figure 1, the results of the study indicated that the dose of adalimumab-based therapy was increased in 9.4\% of patients, while the frequency of dose was increased in 4.0\% of them. For patients receiving infliximab-based therapy, the dose increased in 10.0\% of the cases and the frequency in 11.3\% of them.

The base case economic analysis is presented in Table 3, whilst Figures 2 and 3 present probabilistic results. The mean annual total cost per new (or per existing) responder patient on etanercept was estimated at €9,845 (€9,840), and the total cost on etanercept/MTX was estimated at €9,857 (€9,852). Therapy with etanercept had lower annual cost relative to its comparators. In particular, on an annual basis, it was estimated that the difference between etanercept monotherapy and adalimumab monotherapy was €544 (€1,323). Similarly, the difference between etanercept/MTX and infliximab/MTX was €1,871 (€1,490) and €543 (€1,323), respectively, relative to adalimumab/MTX.

It must be highlighted that, in accordance with the base case scenario, the results imply that for every 1,000 new patients prescribed with etanercept/MTX relative to infliximab/MTX, there are €1.871 million annual savings, and relative to adalimumab/MTX, there are €0.543 million savings. Also, for every 1,000 patients prescribed with etanercept monotherapy, there are €0.544 million annual savings relative to adalimumab monotherapy.

The results of the probabilistic analysis are reflected in the uncertainty confidence intervals in the aforementioned table. In accordance with the probabilistic results, etanercept represents the option with the lowest cost compared with comparators, indicating a statistically significant difference based on 95\% level of significance. The two scenario analyses are presented in Table 4. In the first scenario analysis, the cost of an existing patient responding to therapy with etanercept was estimated at €10,086, followed by etanercept/MTX (€10,098), infliximab/MTX (€9,133), adalimumab (€10,868), and adalimumab/MTX (€10,880). In the second scenario, the cost for etanercept, etanercept/MTX, infliximab/MTX, adalimumab, and adalimumab/MTX were estimated to be €9,840, €9,852, €10,397, €9,916, and €9,927, respectively.

**Table 2** The initial dosage scheme of infliximab

<table>
<thead>
<tr>
<th>Dosage scheme</th>
<th>% of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 mg/kg every 8 weeks (as per label)</td>
<td>70.20%</td>
</tr>
<tr>
<td>5 mg/kg every 8 weeks</td>
<td>14.30%</td>
</tr>
<tr>
<td>3 mg/kg every 6 weeks</td>
<td>11.90%</td>
</tr>
<tr>
<td>5 mg/kg every 6 weeks</td>
<td>3.60%</td>
</tr>
<tr>
<td>Total</td>
<td>100.00%</td>
</tr>
</tbody>
</table>

**Figure 1** Percent of cases of dose escalation and frequency intensification. **Abbreviation:** MTX, methotrexate.
The present study was to assess the economic impact of use of biologic agents for RA treatment on the local payer budget. Considering that no substantial efficacy and safety differences among agents exist, a cost-minimization analysis was considered as the appropriate methodological approach to evaluate alternative therapies from an economic perspective. Adalimumab, etanercept, and infliximab represent the standards of practice in the management of RA patients in Greece. In this analysis, etanercept, alone or in combination with MTX, represents the least costly option and has lower financial impact in the Greek health care payer relative to adalimumab and infliximab. In particular, based on the present analysis conducted, etanercept represents the least costly option for existing and new patients responding to therapy as well as for those not responding to therapy, followed by adalimumab and then infliximab.

In the present model, the observed differences in costs amongst these agents were a result of dose escalation and treatment intensification differences in real-life practice as detected by a field-based survey. The present analysis has several strengths. To the best of our knowledge, this is the first economic study conducted for anti-TNFs in RA patients in Greece. It is based on a large and representative sample of practicing physicians, which permits generalizations of the results to a nationwide basis. Moreover, data from a local registry were also considered in the analysis, to study alternative scenarios. The present findings are consistent with those from other economic analyses undertaken in other countries, which have showed that etanercept may have lower cost per treated patient per year for the base case scenario (in €)

Table 3

<table>
<thead>
<tr>
<th>New patients</th>
<th>Existing patients**</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Nonresponders</strong></td>
<td><strong>Responders</strong></td>
</tr>
<tr>
<td>Etanercept</td>
<td>2,276 (NA)</td>
</tr>
<tr>
<td>Etanercept + MTX</td>
<td>2,278 (NA)</td>
</tr>
<tr>
<td>Infliximab + MTX</td>
<td>3,171 (NA)</td>
</tr>
<tr>
<td>Adalimumab</td>
<td>2,293 (NA)</td>
</tr>
<tr>
<td>Adalimumab + MTX</td>
<td>2,296 (NA)</td>
</tr>
</tbody>
</table>

Notes: Values are expressed as mean (95% UI), where UI was the lower and upper uncertainty interval from 10,000 Monte Carlo simulations. Minimum improvement criteria for a responder were the “ACR 20” (the definition requires, among other prerequisites, a 20% improvement in both tender and swollen joint counts) previously treated patients who were continuing therapy and who achieved an improvement greater than the minimum criteria of ACR 20.

Abbreviations: ACR, American College of Rheumatologists; MTX, methotrexate; NA, not available; UI, uncertainty interval.

Discussion

Understanding the relative outcomes of alternative therapy strategies for patients with RA is important in order to ensure that patients not only receive effective but also, efficient or otherwise cost-effective care. The aim of the present study was to assess the economic impact of use of biologic agents for RA treatment on the local payer budget. Considering that no substantial efficacy and safety differences among agents exist, a cost-minimization analysis was considered as the appropriate methodological approach to evaluate alternative therapies from an economic perspective. Adalimumab, etanercept, and infliximab represent the standards of practice in the management of RA patients in Greece. In this analysis, etanercept, alone or in combination with MTX, represents the least costly option and has lower financial impact in the Greek health care payer relative to adalimumab and infliximab. In particular, based on the present analysis conducted, etanercept represents the least costly option for existing and new patients responding to therapy as well as for those not responding to therapy, followed by adalimumab and then infliximab.

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Figure 2

Mean total treatment cost per year for new patient.

Note: Results were based on 10,000 Monte Carlo simulations.

Abbreviation: MTX, methotrexate.
Because of the economic importance of the treatments for RA, a large number of economic evaluations have been conducted to assess their cost-effectiveness, and several reviews have been undertaken to summarize what is known in this field. Evidence of the cost-effectiveness of the different therapies has been contradictory in published studies, due to the heterogeneity of the approaches applied and the assumption needed regarding the relative effectiveness differences between therapy options. Hence, many commentators and national agencies argue for the case of cost-minimization analysis as a preferable approach to inform decision making regarding allocation of health care resources for RA patients. This is a more straightforward and less controversial approach and is a strong element of the present study. Moreover, the findings from the local survey concerning the dose escalation and frequency adaptation have been consistent with the findings reported in the related literature. For instance, a retrospective review of medical records from five European countries (DART study) also showed that 9.6% of adalimumab patients and 35% of infliximab patients required dose escalation in the

![Figure 3 Mean total treatment cost per year for existing patient.](https://www.dovepress.com/)

**Note:** Results were based on 10,000 Monte Carlo simulations.

**Abbreviation:** MTX, methotrexate.

### Table 4 Average cost per patient per year for two scenario analysis (in €)

| Scenario 1 Dose escalation: 2.5% (etanercept), 9.6% (adalimumab), and 35% (infliximab) (DART study)
<table>
<thead>
<tr>
<th>New patients</th>
<th>Responders*</th>
<th>Existing patients**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Etanercept</td>
<td>2,276 (NA)</td>
<td>9,938 (9,932–9,944)</td>
</tr>
<tr>
<td>Etanercept + MTX</td>
<td>2,278 (NA)</td>
<td>9,949 (9,943–9,955)</td>
</tr>
<tr>
<td>Infliximab + MTX</td>
<td>3,171 (NA)</td>
<td>10,581 (10,496–10,666)</td>
</tr>
<tr>
<td>Adalimumab</td>
<td>2,293 (NA)</td>
<td>10,278 (10,255–10,302)</td>
</tr>
<tr>
<td>Adalimumab + MTX</td>
<td>2,296 (NA)</td>
<td>10,290 (10,267–10,313)</td>
</tr>
</tbody>
</table>

| Scenario 2 Dose escalation: 0% (etanercept), 0% (adalimumab), and 55% (infliximab) (Hellenic Registry)
<table>
<thead>
<tr>
<th>New patients</th>
<th>Responders*</th>
<th>Existing patients**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Etanercept</td>
<td>2,276 (NA)</td>
<td>9,845 (NA)</td>
</tr>
<tr>
<td>Etanercept + MTX</td>
<td>2,278 (NA)</td>
<td>9,857 (NA)</td>
</tr>
<tr>
<td>Infliximab + MTX</td>
<td>3,171 (NA)</td>
<td>11,238 (11,124–11,349)</td>
</tr>
<tr>
<td>Adalimumab</td>
<td>2,293 (NA)</td>
<td>9,921 (NA)</td>
</tr>
<tr>
<td>Adalimumab + MTX</td>
<td>2,296 (NA)</td>
<td>9,932 (NA)</td>
</tr>
</tbody>
</table>

**Notes:** Values are expressed as mean (95% UI), where UI was the lower and upper uncertainty interval from 10,000 Monte Carlo simulations. *Minimum improvement criteria for a responder were the “ACR 20” (the definition requires, among other prerequisites, a 20% improvement in both tender and swollen joint counts). **previously treated patients who were continuing therapy and who achieved an improvement greater than the minimum criteria of ACR 20.

**Abbreviations:** ACR, American College of Rheumatologists; MTX, methotrexate; NA, not available; UI, uncertainty interval.
course of biologic treatment. In addition, another study estimated that 12.5% of adalimumab patients and 42.5% of infliximab patients also underwent dose intensification. Other studies using different methodological approaches showed similar results, consistent with the present findings, which are somewhat lower and hence represent a conservative approach to the issue in hand. Notably, most of the previous analyses concluded that the dose escalation of adalimumab and infliximab were higher than those in the present study, resulting in higher relative cost differentials.

Despite the aforementioned strengths, it is acknowledged that common limitations and drawbacks still exist as the analysis was based on many assumptions and data coming from different sources. In particular, the data obtained from interviews might have been subject to participant recall bias. However, field-based research was considered as a reasonable approach for the purposes of this economic evaluation, in the absence of real-life data at the time of design of this study. In addition, the economic results of this model reflect mainly the specific setting, patient management practice, and drug prices. If these change, so may be the case for the conclusions. Also, the results have to be viewed in the context of the underlying model assumptions. Finally, indirect costs and costs to other parties were not taken into consideration.

A broader analysis from a societal perspective and with case-specific rather than physician data could be the scope of a future economic analysis.

Conclusion
It has been reported that in Greece, RA patients may face access barriers to appropriate biologic therapy. Thus, it is deemed necessary for the health care system to ensure that unrestricted and fairly easy access to all patients is guaranteed, especially in the context of restricted available budgets. In addition, under the pressure of the economic and financial turmoil, there is a growing recognition that comprehensive economic assessments are required, in order to study the economic implications of alternative treatment options and, therefore, to make more informed and better decisions and choices, and to maximize efficiency. The present paper revealed that in the real-life practice setting in Greece, where dose and frequency differences occur, etanercept alone or in combination with MTX, if prescribed as per SPC, represents the option with the lowest annual cost per patient when compared with adalimumab or infliximab in patients with RA. This represents a significant parameter that could be considered in clinical decisions concerning the use of biologic treatments in a cost-constrained environment.

Notwithstanding, the conditions upon which were estimated, these findings may be used by the payer to attain savings in its limited budget.

Author contributions
VF conducted analyses, interpreted the results and drafted the article. EV and ACH designed and supervised study, interpreted results and drafted the article. NM designed study, analyzed and interpreted data and supervised study.

All the authors revised and approved version to be published and are accountable for all aspects of work.

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Disclosure
For the present study, VF has received an unrestricted grant from Pfizer. EV and AH are full time employees at Pfizer Hellas. VF and NM have received honoraria for lectures and seminars from Pfizer. The authors report no other conflicts of interest.

This paper does not guarantee the appropriateness of any medical treatment and is not intended to be used for therapeutic purposes or as a substitute for a health professional’s advice.

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