Economic evaluation of spondyloarthritis: economic impact of diagnostic delay in Italy

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Background and aim: Spondyloarthritis (SpA) is a disease that normally affects the axial skeleton. It progressively leads to overall stiffness up to severe postural deformity of rachis and functional impotence. The objective of the study was to quantify, through an economic model, the impact of specialized testing and pharmacological treatments carried out by the National Health Service (NHS) in normal clinical practice, before the patient is diagnosed with SpA in Italy. In line with the analysis objective, the chosen perspective is that of the NHS.

Method: The study was conducted by analyzing the Health Search Database – IMS Health Longitudinal Patient Database, from which newly diagnosed SpA patients were identified over the period 1 January 2007 to 31 December 2013. The use of specialist health care services and pharmacological treatments provided to the patients before the final SpA diagnosis was estimated.

Results: Through a retrospective analysis of the Health Search Database, 1,084 subjects (aged 25–45 years) were identified. These patients produced an expense of approximately €153,000 in the 3 years prior to a confirmed SpA diagnosis, in terms of specialist check-ups and drugs, presumably not appropriately used due to a lack of diagnosis. If we assume that the Health Search Database is a representative sample of the Italian population, it may be estimated that, in the 3 years prior to SpA diagnosis, over €5.4 million was largely unduly spent in Italy to examine and manage 38,232 newly diagnosed SpA patients, between 2010 and 2013.

Conclusion: The costs due to the delay in SpA diagnosis were quantified for the first time in Italy. For this reason, this work represents a contribution for national and regional decision makers to understand the current clinical practice and the economic consequences of a diagnostic delay in the short and medium term.

Keywords: spondyloarthritis, cost of illness, direct cost, claims database, Italy

Introduction

Spondyloarthritis (SpA) is a disease that normally affects the axial skeleton. It progressively leads to overall stiffness up to severe postural deformity of rachis and functional impotence. SpA also includes psoriatic arthritis, arthritis associated with inflammatory bowel disease, and reactive arthritis. The term SpA covers both patients who have already developed structural damage in the sacroiliac joints or spine visible on radiographs (radiographic axial SpA, also termed ankylosing spondylitis [AS]) and patients without such structural damage, labeled as non-radiographic SpA. The disease usually starts in the third decade of life with a male to female ratio of two to one for radiographic SpA and of one to one for non-radiographic SpA. More than 90% heritability...
has been estimated, the highest genetic association being with HLA-B27. Patients present with chronic back pain and stiffness predominantly of the pelvis and the lower back, but any part of the spine can be involved. A typical symptom is inflammatory back pain, which is clinically defined. Patients complain of morning stiffness mostly of the lower back with improvement on exercise but not by rest. They can also be awakened by back pain in the night, typically in the second part of the night. Several clinical indices (presence of chronic back pain starting at an age ≤ 45 years, inflammatory back pain, peripheral and extra-articular manifestations, response of symptoms to non-steroidal anti-inflammatory drugs [NSAIDs], family history of SpA and related disorders), and several laboratory indices (HLA-B27 and acute phase reactants testing, and imaging findings) have to be combined for classification and diagnosis of SpA. The development of new criteria for classification and for screening of patients with axial SpA have been crucial for the early identification and treatment of such patients, with magnetic resonance imaging being the most important existing imaging method.

The reported prevalence of SpA is between 0.32% and 1.4% in different surveys. According to the ethnic group and the diagnostic criteria used, prevalence of AS ranges between 0.06% and 0.2%. Its incidence is approximately 6–7 patients in 100,000 adults per year. This disease causes high use of health services, reduced productivity, and a limited working capacity in the patients.

SpA has a significant impact on the patient’s life, especially in terms of quality of life. Literature data indicate that the patients are subject to different physical constraints, causing limited working capacity and affecting social relationships and family life. From an economic point of view, the main cost factor is represented by the reduced physical functions.

In terms of economic burden, many studies demonstrated that SpA (mainly AS) causes high social and medical costs. Indirect costs linked to the days of absence from work – together with the loss of related productivity – are the most important determinant of total costs.

This can partially explain the association between total costs and disease severity. A recent revision by Palli et al summarizes the estimated direct and indirect costs in different countries.

As of today, the economic burden of SpA in Italy is not directly known, but some studies estimated that the annual social cost incurred for AS patients treated with biological drugs accounts for approximately €20 million in Italy. These patients represent only 22% of the diagnosed ones.

Diagnostic delay is one of the key clinical challenges of SpA. The average time span from the symptom onset to the definitive diagnosis is 8–10 years. This diagnostic delay has a negative impact not only on the patient, but potentially, also on the disease progression and on the directly related costs incurred by National Health Service (NHS) and society as a whole (indirect costs). The delayed diagnosis causes delay in management and in potential advantages derived from the slowdown of the disability and the improvement of the quality of life.

A delay in diagnosis of 8.57 years was estimated in the UK population, causing loss of spine mobility and inability to work in working-age patients. NSAIDs and blockers of cytokines TNF-α and IL-17 are effective therapies. Early treatment with NSAIDs, as well as a diagnosis within 4 years from the disease onset in patients starting an anti-TNF therapy, may reduce the progression of the radiographic damage. Anti-TNF therapy gives a better response if the duration of the disease symptoms is shorter than 10 years. The main reason for the delay in diagnosis is due to the fact that spinal pain is an extremely common condition and, in most cases, it is not due to severe causes. Furthermore, old diagnostic criteria require significant radiographic lesions, normally arising at a later stage, while routine laboratory tests often supply results in the range for this kind of disease. Conversely, the new diagnostic criteria available today should allow an earlier diagnosis of SpA patients.

The objective of the study was to quantify, through an economic model, the impact of specialized testing and pharmacological treatments carried out by the NHS in normal clinical practice, before the patient is diagnosed with SpA in Italy. In line with the analysis objective, the chosen perspective is that of the NHS.

**Methods**

The study was conducted by analyzing the HSD, from which newly diagnosed SpA patients were identified over the period 1 January 2007 to 31 December 2013. The use of specialist health care services and pharmacological treatments provided to the patients before the final SpA diagnosis were estimated.

**Data sources**

Data were processed through a descriptive inquiry of the HSD, created by the Italian Society of General Medicine and Primary Care in 1998. The database includes a wide and representative sample of the Italian population and is aimed at collecting information derived from the daily clinical practice of general practitioners (GPs).
GPs are selected so as to include a proportional number of patients to the population size of the respective geographical areas. When this study was conducted, Health Search contained information of over 900 Italian practitioners, and a total sample of more than 1.5 million patients.

According to the selection criteria used by the Health Search network researchers, the population represented in the database is distributed by sex and age, in line with the general Italian adult population of ISTAT data.21 The information collected by each GP is representative of the daily clinical practice and contains demographic and clinical information (i.e., diagnoses, laboratory tests with results, hospitalizations, etc.), drug prescriptions, prevention data (i.e., blood pressure, body mass index, smoking, etc.). These data are associated with an identification code of the patient, according to the classical criteria of a relational database.

Furthermore, a lot of this information is directly codified according to the standards commonly used by the NHS for reimbursements, applying the national health care range of fees. For each single drug a pharmaceutical database reports the Anatomical Therapeutic Chemical classification (ATC code), the molecule, the trade name, and the MINSAN code (the code assigned by the Health Ministry to each specific drug when placed on the market).

Enrollment criteria
First, the population to be included in the economic analysis was identified. Specifically, within the database, the patients with at least an SpA diagnosis were identified (code ICD-9 720: “Ankylosing spondylitis and other inflammatory spondylopathies”) over the period 1 January 2010 to 31 December 2013. This code includes the following subgroups of diseases:

- 720.0 “Ankylosing spondylitis”;
- 720.1 “Spinal enthesopathy”;
- 720.2 “Unclassified sacro-iliitis”;
- 720.8 “Other inflammatory spondylopathies”;
- 720.9 “Unspecified inflammatory spondylopathy”.

Among the SpA-diagnosed subjects, only the patients who had received a confirmed diagnosis (by the GP with 720 diagnosis code) during the following year were selected. Finally, in order to identify only the newly diagnosed patients, the subjects who had used health care services due to SpA in the previous 3 years were excluded.

Therefore, all the newly diagnosed SpA patients during the year were selected.

The economic analysis was based only on the patients aged 25–45 years, in order to consider an age group that actually included newly diagnosed SpA patients. The selection process of the population being considered is reported in Figure 1.

Economic analysis
Based on the selected population, all specialist and pharmaceutical services correlating with potential symptoms or manifestations of SpA that took place in the 3 years prior to the first SpA diagnosis were considered, in order to quantify the amount and type of health care services provided before the real diagnosis. In particular, according to the clinical expert opinion of two rheumatologists, co-authors of this study, the SpA-specific health care services that could have allowed an early diagnosis of SpA (Supplementary material) were selected. In order to have a conservative analysis, only services strictly related to SpA diagnosis were considered.

The economic quantification of specialist services was carried out matching the reimbursement fee of the national health care range of fees22 to each service and adding up the costs incurred by each patient in the period of analysis.

![Figure 1 Selection criteria for new SpA patients aged 25–45 years.](https://www.dovepress.com/)

**Abbreviation:** SpA, spondyloarthritis.

Figure 1 Selection criteria for new SpA patients aged 25–45 years.

Period of analysis observation

1 January 2007 → No diagnosis in the previous 3 years

1 January 2010 → Confirmed diagnosis in the following year

31 December 2013

Cost determination → Enrollment period → Follow-up

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The same procedure was carried out for pharmaceutical prescriptions, taking into account the prices available in the pharmaceutical manual associated with each AIC code being analyzed (code of marketing authorization issued by the Italian Medicines Agency).23

In order to obtain a reliable quantification of the expense incurred for these services nationally, the economic model assumed that the analyzed population is a representative sample of the national population. Therefore, an inference analysis was carried out, adjusting the services and related costs to the total resident population in Italy in the same period of analysis.

**Ethical statement**

Ethical approval was not required due to the Health Search IMS Health Longitudinal Patient Database (HSD) being a longitudinal primary care medical record database that was set up by members of the Italian College of General Practitioners (SIMG). The HSD contains anonymized clinical data (diagnoses, patient referrals, hospital admissions, and clinical investigations’ results) and prescription data (drug name, prescription date, number of days’ supply given) for all the medications which are reimbursed by the Italian NHS, and is available to researchers on request.

**Results**

In the total population of the Health Search sample, 6,461 in 1,549,417 subjects received at least an SpA diagnosis over the considered period of analysis. Of these, 3,142 patients (48.6%) were excluded due to the lack of diagnosis by the GP in the following year. An additional 958 patients were excluded because they were already diagnosed as having SpA in the 3 years prior to the index date. Finally, the patients’ extraction in the reference age group was 1,084 subjects, corresponding to an incidence of 2.2 per 1,000 in the 25–45 year old population of the analyzed sample.

Figure 2 shows the method used to identify the objective population, while Table 1 indicates the characteristics of the extracted subjects.

The drug consumption of the 1,084 considered patients during the 3 years of survey amounts to €27,260 in total (Table 2). The most used drugs are glucocorticoids, propionic and acetic acid derivatives, and related substances. The number of specialist services accounts for 4,005 in total.

![Figure 2 Selection process of new SpA patients.](https://www.dovepress.com/)

*Abbreviations: ICD-9, International Statistical Classification of the Diseases and Related Health Problems, version 9; SpA, spondyloarthritis.*
More than half of these correspond to specialist check-ups and laboratory tests, with a total expense of €125,507.

More broadly, it is estimated that in the 3 years prior to an SpA diagnosis, each patient is provided with four specialist health care services and four boxes of drugs related to undiagnosed SpA, with a total expense of €140.9.

Assuming that the Health Search database provides a representative sample of the Italian population and an inference analysis can be carried out, it may be estimated that in Italy, in the 3 years prior to an SpA diagnosis, an expense of over €5.4 million was incurred to examine and treat 38,232 new patients between 2010 and 2013 (Table 3).

**Discussion**

SpA is a disabling disease preventing patients from living a normal life. Early diagnosis and adequate treatment may slow the disease progression down and improve the patient’s health conditions and quality of life in the short and long term. The timespan between the disease onset and the diagnosis is mainly due to difficult early diagnosis of SpA.17,18

For the first time in Italy, this work attempted to quantify the impact of the diagnostic delay, both in terms of services provided and costs incurred by the NHS. Through a retrospective analysis of the Health Search database, 1,084 subjects aged 25–45 years were identified. These patients produced an expense of approximately €153,000 in the 3 years prior to a confirmed SpA diagnosis, in terms of specialist check-ups and drugs, presumably not appropriately used due to a missing diagnosis. According to our assumptions, this cost represents the economic impact produced by the missed SpA diagnoses that might have been reduced if a correct and early SpA diagnostic procedure had been applied.

### Table 1 Characteristics of enrolled patients aged 25–45 years and related diagnoses

<table>
<thead>
<tr>
<th>Clinical characteristics</th>
<th>Patients (n)</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong> (mean)</td>
<td>43.32</td>
<td></td>
</tr>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>381</td>
<td>35.1%</td>
</tr>
<tr>
<td>Female</td>
<td>703</td>
<td>64.9%</td>
</tr>
<tr>
<td><strong>Geographic distribution</strong>*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>North East</td>
<td>313</td>
<td>28.9%</td>
</tr>
<tr>
<td>North West</td>
<td>290</td>
<td>26.8%</td>
</tr>
<tr>
<td>Centre</td>
<td>161</td>
<td>14.9%</td>
</tr>
<tr>
<td>South – Islands</td>
<td>290</td>
<td>26.8%</td>
</tr>
<tr>
<td><strong>Charlson Index</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–1</td>
<td>983</td>
<td>90.7%</td>
</tr>
<tr>
<td>2–3</td>
<td>91</td>
<td>8.4%</td>
</tr>
<tr>
<td>≥4</td>
<td>10</td>
<td>0.9%</td>
</tr>
<tr>
<td><strong>SpA-related diagnoses</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Peripheral enthesopathy and allied syndromes</td>
<td>248</td>
<td>12.4%</td>
</tr>
<tr>
<td>Psoriasis and related conditions</td>
<td>56</td>
<td>2.8%</td>
</tr>
<tr>
<td>Regional enteritis, Crohn’s disease, granulomatous enteritis</td>
<td>11</td>
<td>0.6%</td>
</tr>
<tr>
<td>Ulcerative colitis</td>
<td>9</td>
<td>0.5%</td>
</tr>
</tbody>
</table>

*Note: *The geographic origin of 30 patients is missing. 
Abbreviation: SpA, spondyloarthritis.

### Table 2 Drug and specialist service expense of 1,084 patients aged 25–45 years related to the 3 years prior to first diagnosis

<table>
<thead>
<tr>
<th>ATC code</th>
<th>Drug description</th>
<th>Patients</th>
<th>Drug box consumption</th>
<th>Total expense</th>
</tr>
</thead>
<tbody>
<tr>
<td>M01AH</td>
<td>Coxib</td>
<td>164</td>
<td>389</td>
<td>€6,010</td>
</tr>
<tr>
<td>H02AB</td>
<td>Glucocorticoids</td>
<td>581</td>
<td>1377</td>
<td>€4,873</td>
</tr>
<tr>
<td>M01AE</td>
<td>Propionic acid derivatives</td>
<td>578</td>
<td>968</td>
<td>€4,722</td>
</tr>
<tr>
<td>M03BX</td>
<td>Other centrally acting muscle relaxant drugs</td>
<td>240</td>
<td>310</td>
<td>€3,799</td>
</tr>
<tr>
<td>M01AB</td>
<td>Acetic acid derivatives and related substances</td>
<td>485</td>
<td>919</td>
<td>€3,471</td>
</tr>
<tr>
<td>M02AA</td>
<td>Topical non-steroidal anti-inflammatories</td>
<td>131</td>
<td>149</td>
<td>€1,762</td>
</tr>
<tr>
<td>M01AX</td>
<td>Other non-steroidal anti-inflammatory/antirheumatic drugs</td>
<td>202</td>
<td>511</td>
<td>€1,733</td>
</tr>
<tr>
<td>M01AC</td>
<td>Oxicam derivatives</td>
<td>99</td>
<td>182</td>
<td>€860</td>
</tr>
<tr>
<td>H02BX</td>
<td>Corticosteroids for systemic use</td>
<td>5</td>
<td>7</td>
<td>€31</td>
</tr>
</tbody>
</table>

**Specialist service Patients Number of services Total expense**

| Magnetic resonance diagnostic | 263 | 356 | €41,971 |
| Specialist check-up | 1,012 | 1,700 | €34,445 |
| Laboratory | 493 | 1,095 | €16,293 |
| X-ray diagnostic | 534 | 629 | €14,230 |
| Related diagnostic | 86 | 123 | €9,823 |
| Scintigraphy diagnostic | 28 | 36 | €7,126 |
| Ultrasound diagnostic | 47 | 57 | €1,532 |
| Physiotherapy | 7 | 9 | €87 |
| Total | – | 4,005 | €125,507 |

**Total expense** €152,767

### Table 3 Inferential analysis on the Italian resident population aged 25–45 years between 2010 and 2012

<table>
<thead>
<tr>
<th>Data referred to the analysis</th>
<th>General population (aged 25–45 years)</th>
<th>SpA patients</th>
<th>Cost of services and drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>HS sample</td>
<td>500,842</td>
<td>1,084</td>
<td>€152,767</td>
</tr>
<tr>
<td>Inference on the Italian population</td>
<td>17,664,275</td>
<td>38,232</td>
<td>€5,387,972</td>
</tr>
</tbody>
</table>

*Note: Mean of resident Italian population 2010–2014. 
Abbreviation: SpA, spondyloarthritis.*
If we assume that the Health Search database is a representative sample of the Italian population, it may be estimated that, in the 3 years prior to the SpA diagnosis, over €5.4 million were unduly spent in Italy to examine and manage 38,232 newly diagnosed SpA patients, between 2010 and 2013. The GP plays a key role and should be urged to promptly investigate suspected SpA-related back pain. In specific cases – patient’s young age, inflammatory pain, often at night, alternating sciatica, family history, and concomitant diseases like psoriasis, uveitis, dactylitis, enthesitis – the GP should refer the patient to a rheumatologist.

Like all retrospective analyses on sample data in this field, this study has methodological limits. The first limit is the nature of the Health Search database. In fact, from a general medicine standpoint, it is possible that the diagnosis is not reported on time or the patient’s diagnostic workup is not completed by the GP but by a specialist. In fact, of the 6,461 patients with at least an SpA diagnosis in the analyzed period, approximately 48% were not confirmed the diagnosis the following year. This may be due to a false diagnosis or to the fact that the patient is followed by a specialist and the GP has lost track of the patient. In both cases this study underestimates the real impact of the health expense.

The second limit to the analysis is linked to the identification of SpA patients and to the inclusion criteria. Specifically, the diagnosis code considered in the analysis was ICD-9 720 “Ankylosing spondylitis and other inflammatory spondylopathies.” In effect, this code represents a generic value that may also include diagnoses not specifically related to SpA, such as spinal enthesopathy (720.1), unclassified sacroiliitis (720.2) or other inflammatory spondylopathies (720.8). However, if “ankylosing spondylitis” had been associated only with code 720.2, it would have caused an extreme loss of information due to the missed compilation of the complete code by the GP. In fact, over 80% of the ICD 9 codes considered did not have a complete codification up to the fourth digit. This also causes incidence estimates to be much higher than reported in the literature.

Finally, it should be considered that the expense obtained by the analysis is underestimated, as it only includes the direct costs for specialist and pharmaceutical services. These costs, calculated at rate, should be added to those related to disabilities and irreversible lesions due to diagnostic delay, in addition to the costs due to the loss of productivity.

In conclusion, the costs due to the delay in SpA diagnosis were quantified for the first time in Italy. For this reason, this work represents a contribution for national and regional decision makers to understand the current clinical practice and economic consequences of a diagnostic delay in the short and medium term.

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Disclosure
The authors report no conflicts of interest in this work.

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