Evidence-based process for decision-making in the analysis of legal demands for medicines in Brazil

Processo de tomada de decisão baseado em evidências na análise das demandas judiciais de medicamentos no Brasil

Proceso de toma de decisiones, basado en evidencias científicas, desde el análisis de demandas judiciales de medicamentos en Brasil

Abstract

Legal actions have been playing a significant role as an alternative pathway to access to medicines in Brazil. These lawsuits demand medicines used in Primary Health Care as well as medicines that are still in clinical research and have not been market approved by the Brazilian National Agency for Sanitary Surveillance (ANVISA). The goal was to analyze medicines demanded through lawsuits brought to the judicial district which includes the city of Rio de Janeiro, Brazil, from July/2007 to June/2008. The medicines in 281 lawsuits were examined for their respective indications, classified according to the presence in publicly-funded lists, market approval by ANVISA, compliance with national clinical guidelines, existence of alternative therapies in lists and support of indication by scientific evidence. Six different categories were described, which are deemed useful to managers and the Judiciary in decision-making. The support of evidence is of utmost importance for medicines that are not included in publicly funding lists and also for those with no available therapeutic alternatives.

Evidence Based Medicine; Pharmaceutical Services; Right to Health; Health Technology Evaluation

Resumo


Medicina Baseada em Evidências; Assistência Farmacêutica; Direito à Saúde; Avaliação de Tecnologias de Saúde
Introduction

Brazil’s 1988 Constitution is a legal landmark that defines healthcare as a universal right which must be guaranteed by the Government. The Brazilian Unified National Health System (SUS) is founded on various principles, the most significant being universal access to healthcare and comprehensiveness. SUS has the responsibility of guaranteeing comprehensive therapeutic services, including pharmaceutical services.

Approval by the Brazilian National Agency for Sanitary Surveillance (ANVISA) allows the pharmaceutical product to enter the market in the country; nevertheless market approval does not signify availability or funding by SUS. The incorporation of medications into SUS is carried out by a complex evidence-based process of medicine selection, involving health authorities at federal, state and municipal government levels which results in medicine lists belonging to three different funding components of pharmaceutical services. The first one is the so-called basic component, which refers to medicines used in primary health care. The second component refers to strategic medicines which includes those used to treat endemic conditions, such as infectious diseases (e.g. HIV/AIDS, tuberculosis, Hansen’s disease, leishmaniasis, malaria) and specific programs (e.g. smoking, lupus erythematosus and multiple myeloma). The third component deals with specialized medicines, those used in the high-cost treatment of rare conditions and also for second or third-line treatment options of highly prevalent diseases. For this third component there is an access strategy at the outpatient level, with diagnostic and therapeutic criteria established by Clinical Protocols and Therapeutic Guidelines developed and updated by the Brazilian Ministry of Health (PCDT).

In spite of the progress in pharmaceutical services that has been made over the last decade, there are still unresolved issues surrounding citizens’ access to medicines. Increasingly aware of their rights, citizens have been taking a more active role in seeking access, often resorting to individual litigation. In the last few years there has been an overwhelming amount of legal demands for medicines, many of which involve medicines which have not yet been approved by ANVISA and high-cost medicines, especially those present in the specialized component.

Health technology assessment (HTA) and evidence-based medicine are tools that aim to increase the safe use of technologies, including medicines, through diverse and complementary analytical strategies. HTA assists decision-making by supporting the formulation of policies in the health sector and incorporation of technology. HTA usually employs summarizing methodologies (e.g. systematic reviews and meta-analyses) and economic assessments in healthcare (e.g. cost-effectiveness and cost-utility analyses), regarding a specific technology in subsidizing health decisions for a specific population. Evidence-based medicine, on the other hand, uses mostly the same analytical methods, but focuses on the individual. It integrates clinical practice and medical skills with the best assessment of external clinical evidence to improve the quality and effectiveness of interventions in healthcare provided to individual patients. Evidence-based medicine introduces the idea of using the best available evidence organized as a “hierarchy” of evidence, for which quality of the study and of the study design are the key.

The decision to adopt a medicine for use in healthcare should be evidence-based. Sound, cumulative evidence is employed to establish clinical and therapeutic guidelines, which, in turn, should subsidize clinical and managerial decision-making in the health system. The use of legal pathways to obtain access to cutting-edge therapies does not imply that evidence may be disregarded; on the contrary, it may be said that because decision-making in legal disputes for medicines is very complex, the use of scientific evidence is an essential part of the process.

The methods for collecting and summarizing evidence stem from HTA and from evidence-based medicine through different and complementary analytical strategies.

This work analyzed medicines demanded by plaintiffs in the judicial district of the city of Rio de Janeiro, Brazil, considering alternative therapies and scientific evidence, in order to establish categories for the decision-making process in SUS, regarding pharmaceutical services management, and in the Judiciary.

Methodology

We conducted a retrospective cross-sectional study. Data were collected from the following: primary lawsuit records from the Court of the State of Rio de Janeiro (source: http://www.tjrj.jus.br, accessed on 12/Sep/2008) database, and appeals records from two databases belonging to the State Health Secretariat (source: Secretaria de Estado de Saúde e Defesa Civil do Rio de Janeiro).

Criteria for inclusion of lawsuits were: individual legal actions filed from July 2007 to June
2008 against the state of Rio de Janeiro at the Judicial District of the Capital (City of Rio de Janeiro) with supply of medicines as subject matter. A total of 1,263 lawsuits adhered to these criteria. Because an estimated 40% of lawsuits present specialized medicines, a sample was randomly selected to reflect this characteristic with 95% confidence. The final sample included 295 lawsuits. Fourteen lawsuits were excluded from the sample after examination of the records, because they had been misfiled regarding the subject matter.

The medicines represented in the resulting 281 actions were checked for approval within ANVISA, as well as for their presence in SUS funding components lists. Indications for medicines listed in the PCDT were additionally checked for compliance with the official protocol. This cross-checking eventually permitted the grouping of medicines with similar status.

Search for supporting evidence was conducted for unlisted medicines and for those prescribed in disagreement with government guideline (possible off label use). The Thomson Micromedex – DRUGDEX System database (http://www.micromedexsolutions.com/micromedex2/librarian, accessed on 20/Jan/2011), was elected as the summarized evidence source because of its widespread availability in the public sector, favoring its access by personnel in health management and in the Judicial System. The scale for strength of evidence and of recommendation was applied to medicines and their respective indications.

The careful description of the requirements for medicines classification within these groups and the result of the search for supporting evidence were the origins of the analytical, mutually exclusive, categories.

The project was approved by the Ethics in Research Committee of the Sergio Arouca National School of Public Health, Oswaldo Cruz Foundation (CEP/ENSP/Fiocruz) under record number 33/09.

Results and discussion

Three hundred and forty-four medicines were represented in the 281 demands. Most of them (229 or 66.6%), were not included in any funding component list and 72 (20.9%) belonged to the specialized component.

These results were not consistent with those published in previous studies that showed that most medicines under litigation were included in public funded lists. 4,5,6,10,11,12. Recent studies have also shown similar results to the ones presented in this paper, which may indicate a change in the lawsuit profile. 13,25,26. This finding may be positive since a change of lawsuit profile may be related to better management of pharmaceutical services. A basic requirement for rational use is that listed essential medications are actually available in the system. 27. Nevertheless, other more down-to-earth explanations must be considered, such as pressure from the pharmaceutical industry on prescribers, regarding newer and costlier medicines, which results in non-adherence of healthcare professionals to the evidence-based funding components lists, a common enough situation in many settings. 4,6,11.

The original grouping of medicines and search for evidence resulted in six mutually exclusive categories of medicines.

The first category consisted of medicines which are included in public funding lists and that presented indications compliant with ANVISA and with national guidelines (PCDT). Examples in this category were pegulated interferon + ribavirin for chronic viral hepatitis C (VHC) and infliximab for rheumatoid arthritis.

As these are medicines that should at all times be available in the health system, demands may reflect failings in pharmaceutical services management. Because the judicial pathway and not an administrative one was followed, it is difficult to ascertain if the patient accessed pharmaceutical services in SUS at any given time and if so, what went wrong. The difficulty to gain access to information regarding the availability of certain medicines by patients and prescribers in SUS has been well established. They may be present in one health unit but not in others, confusing patients and care providers. 28. It is the system’s responsibility to ensure that healthcare professionals and users may access medicines. This should be done by updating information on availability within the healthcare network.

The second category comprises specialized component medicines with indication supported by evidence and by ANVISA’s market approval indication but in disagreement with PCDT therapeutic guidelines. Examples are simvastatin for the prevention of cardiovascular events in diabetic patients and the combination of formoterol and budesonide to treat chronic obstructive pulmonary disease. This category is related to a lack of updated guidelines, or lack of any guidelines, and to failings in the relationship between different bodies. One solution to these conflicts may reside in drawing up updated PCDT. The first PCDT were published in 2002 and updating of existent PCDT and the development of additional guidelines occurred mostly between 2009 and 2010, which demonstrates considerable lag time.
between the two. This interval is unacceptable, in face of the availability of newer, fresher evidence concerning the indications for dispensing specialized component medicines.

Policy implementation experiences in the health sector shows that policy adoption is not always homogeneous. Many constraints may arise from this and the population is sure to be affected. Despite the fact that regulatory activities should be conducted apart from policy and provision in the health system, ANVISA is part of the Brazilian Ministry of Health and both should adhere to the same guidelines. If these government services cannot find agreement on issues regarding indication of publicly-funded medicines, it seems difficult to envision flawless transit of patients and prescribers through the system. Many country experiences show that government accountability at this level is not only possible but essential for dissemination of information to health system users and health care professionals. It is a tool to ensure patient rights as to the best possible available treatment.

In the PCDT for dyslipidemia, simvastatin is indicated for patients who have a high risk of developing cardiovascular events, with lipoprotein metabolism disorders and other dyslipidemia-related diagnoses (CID E.78). There are three criteria for a patient’s inclusion in a statin-based treatment: LDL cholesterol levels above 160mg/dL; acute myocardial infarction and LDL cholesterol levels above 100mg/dL; and patients whose LDL cholesterol levels are above 130mg/dL and have at least one aggravating condition (coronary artery disease, previous myocardial infarction, atherosclerotic disease, diabetes, genetic syndromes such as familial hypercholesterol-emia and familial combined hyperlipidemia or high absolute Framingham risk score: ≥ 9 scores for men or ≥ 15 scores for women) 30.

Only three out of eighteen litigating patients were able meet these criteria, but all eighteen received simvastatin as result of litigation, showing not only non-adherence to best evidence or to existing therapeutic guidelines by prescribers, but lack of compliance to best-evidence practices by the Judiciary.

Sometimes best evidence conflicted with PCDT. Regarding the evidence present in the existing guideline, litigating patients with diabetes would have to show LDL cholesterol levels above 130mg/dL in order to receive simvastatin. However, other international evidence-supported guidelines accept the use of simvastatin in the prevention of cardiovascular events in high-risk patients due to cardiovascular atherosclerotic disease or diabetes, irrespective of their cholesterol levels.

Off label indications for simvastatin, such as dementia, were present in lawsuits. However, newer evidence, present in a systematic review of double-blind, randomized clinical trials with a total of 26,340 patients concluded that there is no sound evidence to support this off-label indication. Simvastatin was granted, anyway.

The pathological conditions of patients litigating for formoterol and budesonide were non-specified chronic obstructive pulmonary disease, mixed asthma and non-specified asthma. The PCDT for asthma recommends the use of associations of medicines. However, although ANVISA has also approved this association for use in cases of chronic obstructive pulmonary disease based on sound scientific evidence, no PCDT for chronic obstructive pulmonary disease exists, which prevents dispensing.

The third category includes all medicines which are unavailable in the system and not yet incorporated by SUS, and for which no publicly financed therapeutic alternative exists. However, the use of these medicines is evidence-based and they have been approved by ANVISA. The examples are travoprost for glaucoma and ursodeoxycholic acid to treat primary biliary cirrhosis.

Pilocarpine and timolol are used as first-line treatment options for glaucoma and their use is supported by evidence but these medicines have not been funded over the years. It is important to highlight that the alternatives have been included in the National Essential Medicines List (RENAME) since 2006. Timolol is currently present in the essential medicines list for primary health care of the state of Rio de Janeiro.

Travoprost was requested by sufferers from primary open-angle glaucoma or from non-specified glaucoma. Therefore, while no lack of evidence prevents the medicine from being dispensed, the use of travoprost should occur later in treatment, if patients do not respond well or are intolerant to other anti-glaucoma agents, such as pilocarpine and timolol. The lack of more consistent information about the patients in the lawsuits made it difficult to judge the actual need for travoprost use. In these cases, we may hypothesize that a delay in incorporating first-line anti-glaucoma agents into publicly-funded lists was an important determinant for the onset of legal demands.

There is no therapeutic alternative in publicly-funded lists for ursodeoxycholic acid, and many claims are related to the treatment of primary biliary cirrhosis, for which the use of this drug is supported by evidence and in spite of the fact that it is registered by ANVISA.

A fourth category is characterized by medicines with registered indications by ANVISA, sub-
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statiﬁed by evidence, but indicated mainly as second or third-line treatment options and for which alternatives are funded by the system. The following medicines were part of this category: infliximab for ankylosing spondylitis, clopidogrel for myocardial infarction and losartan for hypertension. Because they are not ﬁrst-choice, analyzing patient history is an important step in the decision-making process. Balancing risks and beneﬁts of use are also worth pondering within this category and further studies may be needed to consolidate available evidence for these medicines and respective indications. Once again it is up to the healthcare system management to take steps to increase adherence of prescribers to the publicly-funded lists and to encourage and stimulate prescription of medicines selected through an evidence-based process and standardized by PCDT.

In this category there are ﬁrst-line alternatives available in SUS. One of the alternative to clopidogrel is acetylsalicylic acid which is listed for the Primary Health Care Component. It is a lower-cost option that is effective and well tolerated by most patients. Clopidogrel is used as an option for patients who are intolerant or who have not responded properly to acetylsalicylic acid. It is necessary to review the patient’s history before furnishing it.

Losartan was included in the 2008 RENAME and in Primary Health Care Component in 2010 but it remains as a second-choice treatment in cases of intolerance to widely-used angiotensin-converting enzyme inhibitors such as captopril and enalapril, which have been traditionally included in RENAME and in primary health care lists. Patients who requested losartan suffered from hypertension, hypertensive cardiac disease without heart failure and congestive heart failure. The use of losartan would only be justiﬁed if the patient did not tolerate the ﬁrst-line choices.

The ﬁfth category includes medicines for which long-term use in intended indications is not supported by evidence. Insulin analogs to treat types 1 and 2 diabetes were included in this category. The existence of evidence supporting short/mid-term use does not yet seem to justify the addition of such medicines to publicly-funded lists. Existing evidence does not support adoption by health systems or clarify pathways for national use.

Insulin is used to treat patients with type-1 and type-2 diabetes when they do not respond to treatment with oral hypoglycemians. The Brazilian Ministry of Health currently provides only human NPH and regular insulin. Insulin analogs (lispro, aspart and glargine) come from recombinant DNA technology, whose structural modification leads to changes to its pharmacokinetic characteristics.

A meta-analyses with a total of 8,274 patients in randomized controlled trials showed that efﬁcacy in glycemic control in long-term insulin analogs was identical to regular human insulin, and also found similar episodes of hypoglycemia. However, the study warns that there is no information on late complications such as diabetic retinopathy or nephropathy. The same result was found in the study by Davis, whose claim is the clinical response of insulin analogs, and proves to be the same to the bolus administration of regular human insulin, when the injections are made in 10-15 and 30 minute periods prior to meals. According to Wannmacher, the advantages of insulin lispro and aspart are: (1) reduction of 20 to 30% of hypoglycemia, and (2) modest improvement (0.3% to 0.5%) in glycosylated hemoglobin.

The RENAME 2008 didn’t include insulin aspart, lispro and glargine and justiﬁed the exclusion by evidence from randomized clinical double-blind studies that have proven insuﬃcient to ensure a therapeutic advantage of insulin lispro and insulin aspart compared to regular human insulin on treatment of diabetes mellitus type 1. The Technical Committee and Multidisciplinary Update of the National List of Essential Medicines (COMARE) further argued that for patients with type 2 diabetes mellitus, there are no studies of insulin lispro compared to regular insulin, and there is insuﬃcient evidence with the use of insulin aspart. In addition, the schedule of administration of insulin lispro or aspart immediately before meals was not enough to ensure that additional therapy provides some beneﬁt to the recommended regimen of human insulin. About insulin glargine, COMARE’s report reveals that the reduction in glycosylated hemoglobin caused by this insulin is not followed by a lower incidence of severe hypoglycemia compared with other types of insulin. As such, the technical committee did not recommend the inclusion of any insulin analogs in RENAME 2008, or in subsequent national essential medicine lists or national publicly funded lists.

The duration of action of a particular type of insulin varies considerably from one patient to another and even within the same individual and this variability comes from different subcutaneous absorption rates, changes in diet and exercise (ANVISA. Bulário eletrônico 2009. http://www.anvisa.gov.br/bula/, accessed on 20/ Dec/2009). Thus, for the dispensing of insulin analogs, it is necessary to examine the patient’s history.
The sixth category is a category of “rogue” medicines: those which do not yet have market approval in Brazil or those for which indications are explicitly unapproved by ANVISA. This was the case for sulthiame (prescribed for epilepsy), which was not approved by the regulatory agency. As for the latter, various examples were found: infliximab to treat psoriasis, olanzapine prescribed for global development disorders, clopidogrel for peripheral vascular disease, arterial embolism or thrombosis, and ursodeoxycholic acid to treat granulomatous hepatitis, hepatic fibrosis and neonatal jaundice caused by other hepatocellular lesions (ANVISA. Bulário eletrônico 2009. http://www.anvisa.gov.br/fala_bula/, accessed on 20/Dec/2009). Demands do not seem to be justified, even if therapeutic alternatives are lacking for the above indications. No efficacy or safety may be established for these drugs in these indications and patients may be at risk.

The last two categories are the most sensitive with regard to safety of medicine use. For the second, third, fourth and sixth categories the question to ask is if in such cases there is a real need for the prescribed medicines. However there are some circumstances that justify its use: whether the plaintiff has an absolute contraindication to the use of the existing SUS alternative therapies; if, after use of a particular drug there has been no improvement in the clinical profile; or if there has been a true delay in incorporation in essential medicine lists.

The literature has shown that failure in incorporation has justified lawsuits in the past as has been the case regarding specific indications of adalimumab, etanercept, infliximab, levodopa + benserazide, peginterferon, rigavastamine, mesalazine, simvastatin, riluzole 6,9. However other cases are not supported by evidence which suggests that litigation represents pressure for SUS to adopt such medicines, targeting private interests more than health interests or therapeutic needs.

It may be necessary to analyze actual cases to support the relevance of claims when there is scientific evidence that justifies prescribing for the indication but the medicine is not included in official lists 10,33.

Final considerations

The analysis of legal demands and their support by scientific evidence allowed for the creation of categories of demanded medicines aiming to support decision-making and Pharmaceutical Services management. The categories are important to guide the drafting of new PCDT or to update existing ones. Furthermore they may help any process regarding inclusion of medicines in publicly funded lists or contribute to the necessary protection of patients receiving new and possibly unsafe medicines or subjected to irrational off-label use.

It is noteworthy to point out that the sources of data from legal actions often did not provide any information on other morbidities simultaneously experienced by the plaintiff, or the direct relationship between diagnoses and requests for medicines. This issue could impose limits to the analysis of actual cases when there is more than one prescribed medicine and/or more than one diagnosis. Nonetheless, HTA and evidence-based medicine tools helped to determine situations supported by evidence and those devoid of supporting evidence, clearly defined by objective information available in lawsuits.

The appropriate structuring of SUS pharmaceutical services ensures the supply of medicines selected by the system as a vital strategy for its maintenance and credibility. Identifying shortcomings in pharmaceutical services may provide information to healthcare managers on where to intervene in order to enforce the right to health. Pointing to the existence of alternative, safe and effective therapies within SUS while also alerting the system in respect to “rogue” medicines whose effectiveness and safety prevent recommendation of use regardless of indication are both correct actions that aim to guarantee the right to health. Another way of safeguarding the right to health is the careful in-depth analysis of particular cases, demanding more information about the reasons for prescribing so that a cautious decision might be made, preserving the patient and the system.
Resumen

Las demandas han jugado un papel importante como una forma alternativa de acceso a los medicamentos en Brasil. Estas demandas incluyen los medicamentos utilizados en ensayos primarios, incluso los que continúan en investigación clínica y no están registrados en el país por la agencia de salud nacional (Agencia Nacional de Vigilancia Sanitaria – ANVISA). El objetivo fue analizar los fármacos presentes en las demandas de la región de Río de Janeiro durante el periodo de julio/2007 a junio/2008. Los fármacos presentes en 281 demandas fueron examinados por su indicación terapéutica, clasificados de acuerdo con su presencia en las listas de financiación pública, su aprobación por la ANVISA, que indica el cumplimiento de las directrices clínicas nacionales, la existencia de terapias alternativas y la existencia de evidencias científicas. Se describieron seis categorías diferentes, en nuestra opinión, útiles para los gestores de la salud y la Justicia en el proceso de decisión. La búsqueda de evidencias científicas es importante para los medicamentos que no están incluidos en las listas, y también para los que no tienen alternativas terapéuticas.

Medicina Basada en Evidencia; Servicios Farmacéuticos; Derecho a la Salud; Evaluación de Tecnologías de Salud

Contributors

T. A. Figueiredo contributed to the design of the research work’s central idea, data collection, database creation, data analysis, search for scientific evidence and writing of this paper. C. G. S. Osorio-de-Castro participated in the design of the research work’s central idea and to writing this paper and proofreading its final version. V. L. E. Pepe collaborated on the design of the research work’s central idea, data analysis and writing this paper.

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Conflict of interest

None declared.

References


33. Macedo EL. A importância da análise técnica para a tomada de decisão do fornecimento de medicamentos pela via judicial [Dissertação de Mestrado], Sorocaba: Universidade de Sorocaba; 2010.


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