Studies on biosimilar medications

Estudos de medicamentos biosimilares

Winston Bonetti Yoshida*

Abstract

In Brazil, the registration of new drugs is carried out only when the regulatory agency – Agência Nacional de Vigilância Sanitária (Anvisa) – is completely satisfied with the evidence of their quality, efficacy and safety, presented by a pharmaceutical industry that strives for this registration. With the patent expiration, pharmaceutical companies are attracted to produce biological medicines called biosimilar or biogenerics or simply generics, whose approval may result in reduced treatment costs. But it is necessary that the biosimilar be, at least, equally effective and safe and without contaminants in relation to the original. Recent consensus guidelines aim to establish criteria for efficacy and safety of these medicines. Preclinical studies in vitro and in vivo, the origin of raw materials and clinical studies phase I, II and III are recommended for biosimilar medicine registration in the international market. Low molecular weight heparins are found in this situation. In this review we specifically addressed this type of medicine, which could serve as a benchmark for other biosimilar medicines.

Keywords: Heparin; heparin, low-molecular-weight; drugs, generic; practice guidelines as topic; therapeutic equivalence

Introduction

In Brazil, new drugs registration is only done when the regulatory agency – Agência Nacional de Vigilância Sanitária (Anvisa) – is completely satisfied with the evidence of their quality, efficacy and safety presented by a pharmaceutical industry that applied for the registration. Recent consensus guidelines aim to establish criteria for efficacy and safety of these medicines. Preclinical studies in vitro and in vivo, the origin of raw materials and clinical studies phase I, II and III are recommended for biosimilar medicine registration in the international market. Low molecular weight heparins are found in this situation. In this review we specifically addressed this type of medicine, which could serve as a benchmark for other biosimilar medicines.
Biological medications, referred to as biosimilars, biogenerics or simply generics, have different meanings according to the regulatory agencies. With the expiration of patents, pharmaceutical companies produce copies which approval may result in treatment cost reduction. But the biosimilar should be at least as effective and safe as the original medication, besides not having contaminant agents. Slight biochemical and biological differences may bring significant clinical consequences. In the field of Angiology and Vascular Surgery, the low-molecular-weight heparins (LMWH) are currently in this situation. Recommendations for tests that assure such better characteristics of the biosimilars have been published. This kind of medication will be featured in this review, and may serve as parameter for other biosimilars.

The main concern is the origin of the raw material for the production of heparin, and features such as type of tissue and animal and country where it has been manufactured must be displayed as general information. There are reports of heparin contamination with oversulfated dermatan sulfate that reflected on the entire production of heparins and derived products around the globe and have caused deaths in several countries, including Brazil. In a comparative study on heparins in the Brazilian market, there were some samples contaminated by oversulfated dermatan sulfate, and, even those which were not, did not present the same chemical purity pattern and high specific anticoagulant activity as those of the standard product (Liquemine® - Roche). Up to 3% of natural dermatan sulfate is allowed in the product, but no other glycosaminoglycans or impurities are acceptable.

Comparative studies on structural integrity by magnetic resonance imaging technique, molecular weight assessment by Sephacryl S-400 gel filtration, and anticoagulant potential by anti-Xa, anti-IIa activities, among others, may point out the similarity and purity between different preparations. The biosimilar's information regarding these aspects must also be displayed on the original product's monograph, over and above the variations between shares must be similar to that of the original product. Besides, analyses on internal disaccharide sequences and terminal 2,5-anhydro-D-mannose residues by the method of nitrous acid degradation, as well as 1,6-anhydro glucose or N-sulfated glucosamine by heparin treatment method, are equally important. The content of sulfate and carboxyl groups must be described based on measures of conductivity and potentiometric titration. Low-molecular-weight heparins (LMWH) contain 12-20% of antithrombin-binding chains, and this regard may be compared by AT affinity chromatographic techniques, as well as heparin cofactor II activity.

In-vitro tests on the biosimilar's activity must be repeated and coincide with those of the original. Usually, the inhibition of the factors Xa, IIa and aPTT (activated partial thromboplastin time) are employed for this purpose. LMWHs may interact with platelet factor 4 and generate antibodies that stimulate heparin-induced thrombocytopenia (HIT), which presents important clinical implications. These ligations may be quantified by appropriate in-vitro tests. The protamine's capacity of neutralization must also be assessed in comparison with the original product.

Studies on acute and chronic toxicity in at least two animal species in accordance with good laboratory practice guidelines must also be part of the preclinical evaluations, comparing different dosages of the biosimilar and of the originator product. The anticoagulant potential must be gauged using standardized experimental animal models with deep venous thrombosis (DVT) and arterial thrombosis.

The phase I evaluations in normal volunteers for five to seven days should always be performed. The doses must be conventional for the prevention of venous thromboembolism (VTE), as well as the determinations of aPTT, anti-Xa and anti-IIa activity must be obtained and tests to investigate HIT must be conducted. Subsequent investigations in patients with renal failure must guide dosage schemes in this situation.

At least one double-blind phase III study aiming at prevention of arterial or venous thromboembolism is recommended by the European Medicine Agencies (EMEA). For each situation, at least one study of this type would be necessary, namely VTE prevention in risk situations, DVT and pulmonary embolism treatment, and prevention of coronary events in patients with unstable angina.

On phase III, the biosimilar may be assessed in relation to the original product by statistical studies such as superiority, equivalence and non-inferiority trials (Figure 1). Non-inferiority trials are commonly conducted to compare them. Such study intends to determine whether the similar is at least as effective as the original, or even a little worse, but within a preestablished limit, that is, a variation. If it is better than the original, that is, the results are better beyond this variation, there will be a bonus and the non-inferiority result will be equally suitable. Equivalence studies are more restrict and implicate results that are not better or worse, but rather within the pre-established
variation (Figure 1). The non-inferiority margin (Δ) is based on previous studies about the originator product, preferentially in comparison to a placebo. Due to ethical implications, this kind of information is rare, and comparisons to products which are considered to be reference are more common. In non-inferiority trials, the studied populations and the outcome must be equal to those of the study which provided the Δ.

The sample size must take into account the level of the confidence interval (generally 95%), the risk of type II error (incorrect rejection of a non-inferior treatment) or test power and Δ. This margin must be the smallest value presenting an important clinical effect. It is usually variable even in certain studies. There are methods for the calculation of Δ17. The sample size calculation may be done by software similar to that of the Epidemiology and Statistical Laboratory of Universidade de São Paulo (USP), which is available on the internet, among others18. Unfortunately, the samples of the equivalence and non-inferiority trials are frequently very small19. Moreover, it is necessary to emphasize that eventual withdrawals per group, especially due to failures in the interpretation of exams, must be replaced aiming at the maintenance of the project's statistical power.

Phase III studies must assess the patients who concluded all phases and examinations of the project; they are referred to as per-protocol or on-treatment population. Patients who have taken at least one dose of the treatment, including those who interrupted it by any reason, constitute the intention-to-threat population (ITT), which is essential for evaluating medication safety, because adverse effects may manifest in the beginning of the treatment and be the cause of treatment interruption. Another interesting assessment parameter is the number needed to treat (NNT)20. The effects of treatments are better understood by means of this risk measurement, which is the number of patients who need to be treated with the new or concerned treatment to produce a desirable beneficial effect (for instance, headache relief, death or thrombosis prevention etc.) in comparison with a control. It is defined as the inverse of the absolute risk reduction (ARR), that is, facing the risk of a control outcome (pA) and experimental treatment (pB), the absolute risk is pA−pB. The NNT is 1/(pA−pB). The ideal NNT is 1, representing the occasion in which all patients respond to the new treatment and nobody gets better with the control. The higher the NNT, the less effective the new treatment20. When the outcome is an adverse effect, calculation is conducted in the same way, but it is referred to as number needed to harm (NNH)21.

Another aspect to be emphasized is the way the randomization of groups and the blind assessment is made22. It is recommended that, in randomized clinical trials, the process of randomization be done by means of random numbers generated by a computer in order to avoid selection bias. Any other method could be considered technically imperfect. The evaluation of results must also be done by independent examiners without knowledge about the groups, and it is recommended that the agreement between them be investigated by statistical tests23.

In conclusion, the tests that are necessary to the registration of biosimilars must follow strict international protocols because of the variability of biological products, particularly LMWHs. Preclinical in-vitro and in-vivo studies clearly showing the pharmacological similarity and purity, as well as pharmacodynamics test in animals, must precede clinical trials. They should always follow the adequate sequence and design in terms of sample size, inclusion and exclusion criteria, outcome similar to that of the original study, sample calculation and non-inferiority margin, randomization, patients’ withdrawal and replacement, evaluations of ITT patients16,22, and tests in different clinical situations. ANVISA (acronym in Portuguese for the Brazilian Agency of Sanitary Vigilance) has recently organized the III Forum of Biological Medication Update in 2010, and these aspects were addressed and the procedures for approval of this kind of medicines in Brazil were emphasized23. To sum up, scientific evidence and safety must always be above any interest other than the patients’ well-being.
References


Correspondence: Winston Bonetti Yoshida Departamento de Cirurgia e Ortopedia da Faculdade de Medicina de Botucatu da UNESP CEP 18618-970 – Botucatu, SP E-mail: winston@fmb.unesp.br