

Biosimilars







A Comparative Clinical Efficacy and Safety Assessment is the **Final Stage of Demonstrating Biosimilarity**

- Biosimilars are approved based on the totality of evidence1,2
- . The clinical trial aims to confirm that there are no clinically meaningful differences between a biosimilar and its reference product in a sensitive patient population using a sensitive endpoint1,2
- The trial will also assess safety outcomes^{1,2}
- · Equivalence trials are recommended to confirm biosimilarity1,2



Biosimilar Development^{1,2}

During Equivalence Trial Design, a Step-by-step Approach is Used to Determine Equivalence Margins and Sample Size^{3,4}

Identification of study setting

Sensitive population

Sensitive endpoints

Meta-analysis of historical data for the reference product

Size and variability of treatment effect is estimated

Determination of equivalence margins

Based on estimated treatment effect size; represent the largest difference judged to be clinically acceptable

Sample size



Equivalence margins are determined independently for each proposed biosimilar, following discussion and agreement between regulators and the biosimilar developer.²





There are Two Common Statistical Measures Used to Assess Biosimilarity³

Risk difference (RD)

% of patients reaching endpoint with biosimilar - % reaching endpoint with reference product

• If drugs have the same efficacy, RD=0



Risk ratio (RR)

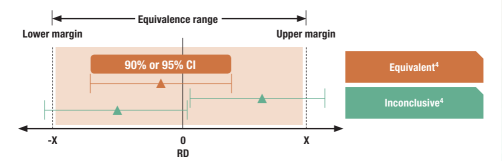
% of patients reaching endpoint with biosimilar

% reaching endpoint with reference product

• If drugs have the same efficacy, RR=1



The Outcomes of RD and RR Analyses are Determined by the Predefined **Equivalence Margins**^{1,2,5}



CI, confidence interval

Comparative clinical efficacy is shown by demonstrating that the two-sided CI for RR or RD falls within the predefined equivalence margins.^{3,4,6}

1. EMA. Guideline on similar biological medicinal products containing biotechnology-derived proteins as active substance: non-clinical and clinical issues, 2014. $A vailable\ at: http://www.ema.europa.eu/docs/en_GB/document_library/Scientific_guideline/2015/01/WC500180219.pdf;\ 2.\ FDA.\ Scientific\ considerations\ in$ demonstrating biosimilarity to a reference product. Guidance for industry, 2015. Available at: http://www.fda.gov; 3. Isakov L, et al. Am J Ther 2016;23:1903–10; 4. Alten R, et al. Semin Arthritis Rheum 2015;44:S2-8; 5. ICH. Topic E 9 statistical principles for clinical trials, 1998. Available at: https://www.ich.org/fileadmin/ Public_Web_Site/ICH_Products/Guidelines/Efficacy/E9/Step4/E9_Guideline.pdf 6. He J, et al. Clin Cancer Res 2016;22:5167-70. Links accessed February 2019



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