





# Biosimilars

## Hot Topic: The Importance of Analytical Characterization in Biosimilar Development



### Biosimilars Are Not Generic Drugs<sup>1</sup>

- Biologics are products produced in genetically-engineered living cells or organisms<sup>1</sup>
- Biosimilars are biologic medicines that are highly similar to the reference product (RP) with no clinically meaningful differences in terms of safety, purity, and potency<sup>1,2</sup>

	Small molecule drugs Including generics 	Biologics Including biosimilars 
Size	<b>Small</b> <sup>3,4</sup>	<b>Much larger</b> <sup>1,3,4</sup>
Structure	<b>Simple</b> and well defined <sup>3,4</sup>	<b>Complex</b> , with many possibilities for post-translational modification <sup>1,3,4</sup>
Manufacturing	Predictable chemical process; <b>identical copies can be made</b> <sup>3</sup>	Manufactured in a unique, living cell line; <b>only similar, not identical copies can be made</b> <sup>3,4</sup>
Characterization	<b>Easy to characterize fully</b> <sup>4</sup>	<b>Difficult to characterize fully</b> <sup>4</sup>
Stability	<b>Relatively stable</b>	<b>Often sensitive to storage and handling conditions</b> <sup>4</sup>
Immunogenicity	<b>Lower potential</b> <sup>4</sup>	<b>Higher potential</b> <sup>4</sup>

### Biosimilar Manufacturers Start with Limited Knowledge of the Reference Product

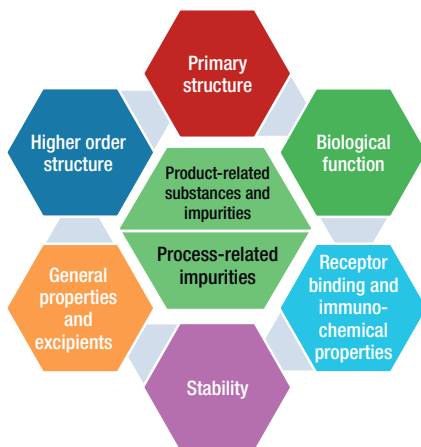
- Thorough characterization of the RP is the first step in biosimilar development<sup>1,2</sup>
- The biosimilar manufacturer must then produce a unique cell line and develop an entirely new manufacturing process that produces a highly similar product<sup>2</sup>



Reference product manufacturing information is proprietary and not publicly available.<sup>2</sup> A biosimilar manufacturer must develop an entirely new customized process.

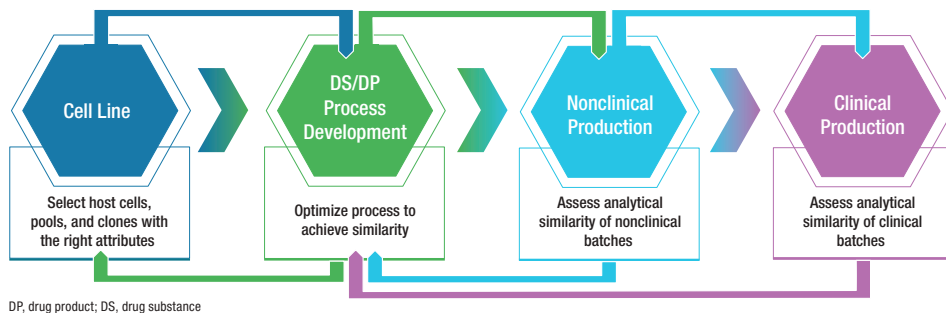
## Analytical Characterization is Used to Evaluate RP Critical Quality Attributes (CQAs) in Eight Categories<sup>5</sup>

- Analytical characterization of the reference product identifies the CQAs<sup>1-4</sup>
- CQAs are specific attributes that impact pharmacokinetics, safety and efficacy<sup>3,4</sup>
- CQAs must be controlled within an appropriate range to ensure product quality<sup>3</sup>



## Biosimilar Development: The Product Defines the Process

- Similarity in structure and function is established via an iterative process<sup>1,2</sup>



- At each stage, the manufacturer evaluates analytical data and determines whether to proceed with development or conduct further optimization

**Analytical similarity assessment is an iterative operation conducted throughout process development.<sup>3,4</sup>**

### References

- FDA. Scientific considerations in demonstrating biosimilarity to a reference product. Guidance for industry, 2015. Available at: <http://www.fda.gov/downloads/>. 2. EMA. Guideline on similar biological medicinal products containing biotechnology-derived proteins as active substance: quality issues (revision 1), 2014. Available at [https://www.ema.europa.eu/documents/scientific-guideline/draft-guideline-similar-biological-medicinal-products-containing-biotechnology-derived-proteins\\_en-0.pdf](https://www.ema.europa.eu/documents/scientific-guideline/draft-guideline-similar-biological-medicinal-products-containing-biotechnology-derived-proteins_en-0.pdf). 3. Markus R, et al. Biodrugs 2017;31:175–87.
- Vulto A, et al. Rheumatology 2017;56:iv14–iv29. 5. FDA. Quality considerations in demonstrating biosimilarity of a therapeutic protein product to a reference product. Guidance for industry 2015. Available at: <http://www.fda.gov/downloads/>. Links accessed November 2018.