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Title	Regulation of biosimilar medicines and current perspectives on interchangeability and policy				
	interchangeability and policy				
Author(s)	O'Callaghan, John; Barry, Sean P.; Bermingham, Margaret; Morris, J. Michael; Griffin, Brendan T.				
Publication date	2018-09-05				
Original citation	O'Callaghan, J., Barry, S. P., Bermingham, M., Morris, J. M. and Griffin, B. T. (2018) 'Regulation of biosimilar medicines and current perspectives on interchangeability and policy', European Journal of Clinical Pharmacology. doi:10.1007/s00228-018-2542-1				
Type of publication	Article (peer-reviewed)				
Link to publisher's	http://dx.doi.org/10.1007/s00228-018-2542-1				
version	Access to the full text of the published version may require a				
	subscription.				
Rights	© 2018, Springer-Verlag GmbH Germany, part of Springer Nature.				
	All rights reserved. The final publication is available at Springer via				
	http://dx.doi.org/10.1007/s00228-018-2542-1				
Embargo information	Access to this article is restricted until 12 months after publication by request of the publisher.				
Embargo lift date	2019-09-05				
Item downloaded	http://hdl.handle.net/10468/7163				
from					

Downloaded on 2019-12-02T14:47:19Z



Electronic Supplementary Material

Regulation of biosimilar medicines and current perspectives on interchangeability and policy

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Journal: European Journal of Clinical Pharmacology

Table S1: Regulatory explanations of the term 'biosimilar'

European Medicines	A biosimilar is a biological medicinal product that contains a version of the					
Agency ¹	active substance of an already authorised original biological medicinal product					
	(reference medicinal product) in the EEA. Similarity to the reference medicinal					
	product in terms of quality characteristics, biological activity, safety and					
	efficacy based on a comprehensive comparability exercise needs to be					
	established.					
U.S. Food & Drug	Biosimilar or biosimilarity means that 'the biological product is highly similar					
Administration ²	to the reference product notwithstanding minor differences in clinically inactive					
	components' and that 'there are no clinically meaningful differences between					
	the biological product and the reference product in terms of the safety, purity,					
	and potency of the product'.					

EEA- European Economic Area

¹ European Medicines Agency (2014) Guideline on similar biological medicinal products

² Section 7002(b) (3) of the Affordable Care Act, adding section 351(i) (2) of the Public Health Services Act

Table S2: Comparability exercise to support major manufacturing process changes - details of clinical studies and extrapolated indications

Product name (Active substance)	Approved indications	Nature of change	Clinical data	Extrapolated indication	Ref
Herceptin® (Trastuzumab)	Early breast cancer Metastatic breast cancer	New formulation for subcutaneous administration	Clinical trials in HER2+ early breast cancer patients	Extrapolation to metastatic setting	[1]
Aranesp® (Darbepoetin alfa)	Anaemia associated with chronic kidney failure Anaemia in adult cancer patients receiving chemotherapy	New master cell bank and new manufacturing technology	Clinical trials in chronic kidney failure patients	Extrapolation to cancer indication	[1]
Avonex® (Interferon beta-1a)	Multiple sclerosis	Changes to master cell bank and manufacturing process after pivotal phase III trials	PK data	MS indication approved without new clinical efficacy trial	[2, 3]

PK; pharmacokinetic, MS; multiple sclerosis

Table S3: Substitution of biological medicines in Australia - Details of reference biological medicines and their biosimilars listed on the Australian Pharmaceutical Benefits Scheme (August 2018)

Substance	Products	Substitution permitted ('a' flag status granted by PBAC)*.	Reason 'a' flag status not granted	Ref
Etanercept	Enbrel® Brenzys®	Yes	N/A	[4]
Epoetin alfa Epoetin lambda	Eprex® Novicrit®	No	At time of authorisation Novicrit has restricted route of administration when compared to Eprex	[4, 5]
Filgrastim	Neupogen® Nivestim®	No	Absence of TGA statement that supports 'a' flagging	[4, 6]
Follitropin Alfa	Gonal-f® Bemfola®	No	Substitution difficult from a practical perspective owing to differences in strengths, number of pens per pack and maximum quantities per brand	[4, 7]
Infliximab	Remicade® Inflectra® Renflexis®	Yes	N/A	
Pegfilgrastim	Neulasta®** Ristempa®**	Yes*	N/A	[4]
Somatropin	Genotropin® Omnitrope®	No	No details provided on PBAC website	[4]

^{*&#}x27;Brand equivalents are accompanied by an 'a' flag on the Pharmaceutical Benefits Schedule. An 'a' flagged medicine may be substituted by a pharmacist at the point of dispensing.

PBAC, Pharmaceutical Benefits Advisory Committee; TGA, Therapeutic Goods Administration

^{**}Neulasta and Ristempa are not biosimilars but are the same products with different brand names

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 Available from: http://www.ema.europa.eu/docs/en_GB/document_library/EPAR_-
 Assessment Report Variation/human/000332/WC500026148.pdf.
- Food and Drug Administration (FDA). Avonex Summary Basis of Approval. 2003 2 Nov 2017]; Available from: https://www.accessdata.fda.gov/drugsatfda_docs/nda/2003/103628_S0000_SBA.pdf.
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- 5. Pharmaceutical Benefits Advisory Committee (PBAC). Public summary document July 2010 Epoetin Lambda injection,1000 units in 0.5ml, 2000 units in 1.0ml, 3000 units in 0.3ml, 4000 units in 0.4ml, 5000 units in 0.5ml, 6000 units in 0.6ml, 8000 units in 0.8ml and 10,000 units in 1.0ml, pre-filled syringe, Novicrit, Novartis Pharmaceuticals Australia Pty Ltd,. 2010 6 November 2017]; Available from:

 http://www.pbs.gov.au/industry/listing/elements/pbac-meetings/psd/2010-07/Epoetin_lambda_NOVICRIT_Novartis2.pdf.
- 6. Pharmaceutical Benefits Advisory Committee (PBAC). Public summary document November 2010 PBAC meeting Filgrastim, injection, 120 micrograms in 0.2ml single use pre-filled syringe, 300 micrograms in 0.5ml single use pre-filled syringe, 480 micrograms in 0.5ml single use pre-filled syringe, Nivestim, Hospira Pty Ltd,. 2010 6 Nov 2017]; Available from: http://www.pbs.gov.au/info/industry/listing/elements/pbac-meetings/psd/2010-11/pbac-psd-filgrastim-nov10.
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