Federal Institute for Vaccines and Biomedicines



Antibody Drug Conjugates as specific and potent Medicinal Products - a regulator's perspective -



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Disclaimer

The view expressed in the following is the ones of the presenter and does not necessary express the view of either the CHMP, BWP, EDQM or the Paul-Ehrlich-Institut (including other sections)

Antibody moiety



high binding affinity to the target antigen, efficient internalization, low Immunogenicity, long plasma half-life



Adapted from Carter PJ: Potent antibody therapeutics by design, Nature Rev Immunology 6, 343 (2006)

- IgG1 is the most abundant in serum and could induce the strong effector functions such as ADCC), ADCP), and CDC
- IgG2 reduced effector function form dimers and aggregations in vivo, which leads to a decrease of the concentration of ADC drugs
- IgG3, short half-life (7 days)
- IgG4 could induce ADCP, reduced effector function

Antibody moiety formats





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From C Siess et al, Alternative molecular formats and therapeutic applications for bispecific antibodies, Molecular Immunology 67 (2015) 95–106

Cleavable vs. non-cleavable linkers



Non-cleavable linkers: low off-target toxicity benefited from an increase of plasma stability (T-DM1)

Cleavable linkers:

- chemical cleavage linkers (hydrazone bond and disulfide bond) Hydrazone is a typical acid-sensitive (pH sensitive) linker.
- enzyme cleavage linkers (glucuronide bond and peptide bond), sensitive to the lysosomal protease e.g. Cathapsin B

Linker Technology:

- PEG/Polysarcosine for solubility/hydrophilic characteristics



Source (Peters and Brown 2015)

Cytotoxic payloads



potent tubulin inhibitors, DNA damaging agents, RNA synthese inhibition and immunomodulator

- Tubulin polymerization promoters target at the β-subunits of tubulin dimer to perturb microtubule growth, auristatin derivatives monomethyl auristatin E (MMAE) and monomethyl auristatin F (MMAF)
- IC50 values of DNA damaging agents are able to reach picomolar level
 - DNA double strand break, such as calicheamicins;
 - DNA alkylation, such as duocarmycins;
 - DNA intercalation, such as topoisomerase I inhibitors (exatecans)
 - DNA crosslink, such as pyrrolobenzodiazepines (PBD).
- Specific inhibition of RNA polymerase II activity
 - α-Amanitin mechanism of action: Cell-cycle independent mechanism of actionLow intracellular target copies, 1:1 binding

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Growing microtubul

Paused microtubule (neither polyme







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Drug-load variants





Conjugation methods



- > stochastic conjugation on pre existing lysine or cysteine residues via appropriate coupling reaction
 - random coupling with lysine residues, varying numbers (0–8) of small-molecule toxins may be attached to an antibody, resulting in a wide drug-antibody ratio (DAR) distribution
 - Cysteine based reaction provides another means of coupling. Due to the limited number of binding sites and the unique reactivity of mercaptan groups, using cysteine as the connecting site helps to reduce the heterogeneity of ADC. Depending on the reduction ratio, products with DAR of 2, 4, 6 and 8 may be generated with better homogeneity compared
- engineered reactive cysteine residues has become a common approach for site-specific conjugation
- Introduction of unnatural amino acids, including N-acetyl-Lphenylalanine, azido methyl-Lphenylalanine and azido lysine, may induce immunogenicity
- from glycan remodeling and glycoconjugation, e.g. N297





ADC development



The first-generation ADCs conventional chemotherapy drug conjugated to a mouse derived antibody through a noncleavable linker. The potency not superior to free cytotoxic drugs immunogenicity was frequently a concern.

The second generation ADCs represented by brentuximab vedotin and ado-trastuzumab emtansine optimization of mAbs isotypes, cytotoxic payloads, as well as linkers.

The third-generation ADCs

The third generation ADC are represented by polatuzumab vedotin, enfortumab vedotin, famtrastuzumab deruxtecan site-specific conjugation technology, well characterized DARs (2 or 4) and desired cytotoxicity were produced, more hydrophilic linker modulation such as PEGylation



Definition starting material

Part II: Basic Requirements for Active Substances used as Starting Materials

An "API Starting Material" is a raw material, intermediate, or an API that is used in the production of an API and that is incorporated as a significant structural fragment into the structure of the API. A Starting Material can be an article of commerce, a material purchased from one or more suppliers under contract or commercial agreement, or produced in house.



EUROPEAN COMMISSION HEALTH AND CONSUMERS DIRECTORATE-GENERAL Health systems and products Medicinal products – guality, safety and efficacy

Volume 4 EU Guidelines for Good Manufacturing Practice for Medicinal Products for Human and Veterinary Use Annex 2: Manufacture of Biological Medicinal Substances and Products for Human, Part B.

Definition and Requirements

"starting materials shall mean any substance of biological origin such as micro-organisms, organs and tissues of either plant or animal origin, cells or fluids (including blood or plasma) of human or animal origin, and biotechnological cell constructs (cell substrates, whether they are recombinant or not, including primary cells)."



London, 27 June 2013 EMA/CHMP/BWP/429241/2013 Committee for Medicinal Products for Human Use (CHMP)

Guideline on the use of starting materials and intermediates collected from different sources in the manufacturing of non-recombinant biological medicinal products

Draft Agreed by Biologics Working Party	December 2011
Adoption by Committee for medicinal products for human use for release for consultation	16 February 2012
End of consultation (deadline for comments)	31 August 2012
Agreed by Biologics Working Party	May 2013
Adoption by Committee for medicinal products for human use	27 June 2013
Date for coming into effect	1 December 2013

Keywords

Starting materials, sourcing, intermediates, heparins, urine derived products, plasma derived medicinal products, manufacturing process.

- The marketing authorisation dossier should include information that acequately describes the manufacturing process and process controls.
- Information on quality and control of all starting materials and process reagents used in the manufacture of a drug substance should be provided.
- GMP measures (e.g. contract between supplier and manufacturer of medicinal product, audit system) should be adequate to ensure an appropriate control while allowing sourcing of starting materials or early intermediate biological products in different locations from third countries.

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ICH Q11, Selection of starting materials and source materials





November 2012 EMA/CHMP/ICH/425213/2011

ICH guideline Q11 on development and manufacture of drug substances (chemical entities and biotechnological/ biological entities)

5. Selection of starting materials and source materials
5.1. General principles
5.1.1. Selection of starting materials for synthetic drug substances
5.1.2. Selection of starting materials for semi-synthetic drug substances
5.1.3. Selection of source and starting materials for biotechnological/ biological drug substances
5.2. Submission of information for starting material or source material
5.2.1. Justification of starting material selection for synthetic drug substances
5.2.2. Justification of starting material selection for semi-synthetic drug substances12
5.2.3. Qualification of source or starting materials for biotechnological/ biological drug substances

5.1.3. Selection of source and starting materials for biotechnological/biological drug substances

Cell banks are the starting point for manufacture of biotechnological drug substances and some biological drug substances. In some regions, these are referred to as source materials; in others, starting materials. Guidance is contained in ICH Q5A, Q5B, and Q5D. Guidance is contained in ICH Q5A, Q5B and Q5D

ICH Topic Q 5 A (R1) Quality of Biotechnological Products: Viral Safety Evaluation of Biotechnology Products Derived from Cell Lines of Human or Animal Origin

ICH Topic Q 5 B Quality of Biotechnological Products: Analysis of the Expression Construct in Cell Lines Used for Production of r-DNA Derived Protein Products

> ICH Topic Q 5 D Quality of Biotechnological Products: Derivation and Characterisation of Cell Substrates Used for Production of Biotechnological/Biological Products





DM1/DM4 Tubulin inhibitor

- Ansamitocins are isolated and purified from the fermentation broth of Actinosynnema pretiosum
- variable product (ansamitocin mixture)
- MayOH is obtained by cleaving the C-3 ester of ansamitocins
- DM1 is synthesized from Maytansinol (MayOH).
- A consistent purity profile of DM1 is achieved by defined and controlled fermentation process.
- Defining the bacteria working seed lot system/fermentation as starting material

Documentation to be provided

- detailed description of the conversion of MayOH to DM1
- Information on the process as well as the in-process controls and/or in-process tests in place
- Information on all reagents and catalyst used in the MayOH conversion to DM1
- Specification, limits and batch data



	•	
Appearance/Description		Visual
Appearance	White to yellow solid	
Identity		
Identity by UV	Conforms	UV
Identity by HPLC	Conforms	HPLC
Purity		
Assay % w/w (dry basis) ^a	93.0% - 103.0%	HPLC
Impurities (area%)		HPLC
Individual specified		
Ring Oxidized MayOH	$\leq 1.0\%$	
O-Desmethyl-MayOH	$\leq 1.0\%$	
Deschloro-MayOH	$\leq 0.50\%$	
N-Desmethyl-MayOH	$\leq 0.50\%$	
Descarbamate-MayOH	$\leq 1.0\%$	
Methyl-MayOH	$\leq 1.0\%$	
AP3	$\leq 1.0\%$	
Individual unspecified impurities	$\leq 0.5\%$	
Total unspecified impurities	$\le 1.5\%$	
Total impurities	$\leq 5.0\%$	
Water Content	$\leq 1.0\%$	KF
Residual Solvents, % w/w		Headspace GC
Ethyl Acetate	$\leq 13.0\%$	
Methanol	$\leq 0.3\%$	
Total Residual Solvents	≤ 15.0%	

Principles of ICH Q11 for classifying starting material for synthetic Drug Substances:



- There is an adequate control strategy in place to control the quality of the proposed starting material.
- There is a reduced risk for fate of impurities due to the number of steps after the introduction of the starting material.
- The DS manufacturing process is sufficiently described to ensure the understanding of formation, fate and purge of impurities.

Starting Material-MMAE



Monomethyl auristatin E is an antimitotic agent which inhibits cell division by blocking the polymerisation of tubulin

MMAE monomethyl auristatin E



convergent, solution phase, fragment-based peptide synthesis.



(Draft) Guideline on the Development and Manufacture of Synthetic Peptides



- new guideline on the development and manufacture of synthetic peptides is currently in preparation
- there is consensus on peptide synthesis aspects, e.g. that in general the protected amino acids are considered regulatory starting materials for synthetic peptides.
- It is still under discussion if bigger building blocks, like several amino acids coupled to a solid phase, may possibly serve as starting material



15 September 2022 EMA/CHMP/QWP/735422/2022 Committee for Medicinal Products for Human Use (CHMP) Committee for Veterinary Medicinal Products (CVMP)

Concept Paper on the Establishment of a Guideline on the Development and Manufacture of Synthetic Peptides

Agreed by Quality Working Party	29 June 2022
Adopted by CHMP for release for consultation	15 September 2022
Adopted by CVMP for release for consultation	8 September 2022
Start of public consultation	20 September 2022
End of consultation (deadline for comments)	20 December 2022

Publishing of the Draft "Guideline on the Development and Manufacture of Synthetic Peptides" is expected end of 2023.

Drug linker commercial specification

Tests for description, identification, related substances, residual solvents and assay are in line with ICH Q6A

- Justification for absence of other physicochemical tests, such as polymorphism and particle size may require further
- The stereochemistry is controlled by material specifications, in-process controls and a test for specific optical rotation in the specification of MAAA-1162a. This control strategy is acceptable when thorough discussion on possible stereoisomers and their fate is provided and supported with analytical data.
- absence of racemisation to be shown with batch analysis data and stability studies.
- risk assessment according to ICH guideline Q3D (R1) on elemental impurities
- ICH guideline Q3C (R7) on impurities: guideline for residual solvents

Description (Appearance)	
Identification	
Specific Optical Rotation (20°C, sodium D line)	
Related Substances Specified – NHS-adduct Individual Unspecified Total	
Residual Solvents 1-Propanol Acetone Ethyl Acetate Tetrahydrofuran (THF)	



Quality guidance for <u>chemical entities</u> applied to the drug-linker intermediate component?

- Drug linker represents an intermediate in the manufacture of the actual DS and might therefore be treated as surrogate API
- It is agreed that for the drug-linker component manufactured using small molecule chemical synthesis, the relevant chemical entity guidance can be applied.
- It is also acceptable to follow Section B.I. Active Substance of the Variation Guidance for "chemical entity drug substance intermediate" for changes concerning the druglinker intermediate only

Caveat:

no DMF (Drug Master File in EU): the concept of the ASMF shall only apply to a welldefined active substance and cannot be used for excipients, finished products and biological active substances

Characterisation/Specifications





Nature Biotechnology 22, 1383 - 1391 (2004)

Maytansinol

Antibody Specifications

Q6B (and ICH Q3D, risk assessment)

- MayOH is a well-characterized, stable compound
- MayOH is a well-defined single compound that is produced and tested for conformance against an appropriate specification using qualified reference standards.

These specifications have to be established based on principles outlined in ICH

MayOH has a well-defined and consistent impurity profile ICH Q6B

MMAE monomethyl auristatin E Specifications have been established based on principles outlined in ICH Q6A Specifications: Test Procedures and Acceptance Criteria for New Drug Substances and New Drug Products: Chemical Substances Specifications.

The specifications set for linker and cytotoxic drug should include the recommend acceptable amounts for residual solvents guidance given in the ICH Q3C "Impurities: Guideline for residual solvents" and ICH Q3D "Elemental impurities" should be followed.



(derivative of maytansine)



MMAE R₁=CH₃, R₂=OH; MMAF R₁=COOH, R₂=H

Need for Guidance on ADCs?

- Clear terminology identifying starting materials, intermediates, linkers, active substance and finished product stages
- Structure of CTD quality and non-clinical modules for intermediates, active substance and finished product
- Reference to the dossier of an already authorised medicinal product (e.g., monoclonal antibodies, radionuclide intermediates) and use of an ASMF procedure for radiopharmaceutical precursors
- Specification requirements for radionuclide, e.g., radionuclide characteristics, radionuclide concentration, radionuclide purity, radiochemical purity, specific activity, chemical composition, chemical impurities, chemical stability
- State-of-the-art radiolabelling method (to generate stable conjugate) requirements
- Specification requirements for active substance and finished product, e.g., identity, purity, potency, sterility





20 July 2023 EMA/CHMP/BWP/245588/2023 Committee for Medicinal Products for Human Use (CHMP)

Concept paper on the revision of the Guideline on Radiopharmaceuticals Based on Monoclonal Antibodies

Agreed by Biologics Working Party	12 July 2023
Adopted by CHMP for release for consultation	20 July 2023
Start of public consultation	21 July 2023
End of consultation (deadline for comments)	31 October 2023



Thank You!



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