



European Medicines Agency

London, 21 May 2008
Doc. Ref. EMEA/CHMP/410869/2006

COMMITTEE FOR MEDICINAL PRODUCT FOR HUMAN USE (CHMP)

GUIDELINE ON HUMAN CELL-BASED MEDICINAL PRODUCTS

DRAFT AGREED BY CPWP AND BWP	November-December 2006
ADOPTION BY CHMP FOR RELEASE FOR CONSULTATION	25 January 2007
END OF CONSULTATION (DEADLINE FOR COMMENTS)	31 July 2007
AGREED BY CPWP and BWP	April 2008
ADOPTION BY CHMP	30 May 2008
DATE FOR COMING INTO EFFECT	1 September 2008

KEYWORDS	Human cell-based medicinal products, quality and manufacturing aspects, Non-clinical development, Clinical development
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TABLE OF CONTENTS

EXECUTIVE SUMMARY	3
1. INTRODUCTION (BACKGROUND)	3
2. SCOPE.....	3
3. LEGAL BASIS	4
4. MAIN GUIDELINE TEXT	4
4.1 RISK ANALYSIS	4
4.2 QUALITY AND MANUFACTURING ASPECTS.....	5
4.2.1 Starting and raw materials.....	5
4.2.2 Manufacturing process.....	8
4.2.3 Characterisation.....	10
4.2.4 Quality control	13
4.2.5 Validation of the manufacturing process	14
4.2.6 Development Pharmaceuticals	15
4.2.7 Traceability	17
4.2.8 Comparability.....	17
4.3 NON-CLINICAL DEVELOPMENT	18
4.3.1. Pharmacology	18
4.3.2. Toxicology	19
4.4 CLINICAL DEVELOPMENT	21
4.4.1 General aspects	21
4.4.2 Pharmacodynamics	21
4.4.3 Pharmacokinetics	21
4.4.4 Dose finding studies	21
4.4.5 Clinical Efficacy.....	22
4.4.6 Clinical Safety	22
4.4.7 Pharmacovigilance and Risk Management Plan.....	23
REFERENCES (SCIENTIFIC AND / OR LEGAL)	23

EXECUTIVE SUMMARY

This guideline replaces the Points to Consider on the Manufacture and Quality Control of Human Somatic Cell Therapy Medicinal Products ([CPMP/BWP/41450/98](#)). It takes into account the current legislation and the heterogeneity of human cell-based products, including combination products. A risk analysis approach can be used by the applicants to justify the development and evaluation plans and can be a basis for the preparation of a risk management plan.

In the quality and manufacturing section, guidance is provided on the criteria and testing of all starting materials, on the design and validation of the manufacturing process, on characterisation of the human cell-based medicinal products, on quality control aspects, on the development programme, traceability and vigilance* and on comparability issues. Guidance specific to the matrix/device/scaffold component in combination products is provided.

The guideline acknowledges that conventional non-clinical pharmacology and toxicology studies may not be appropriate for cell-based medicinal products. Therefore the guideline addresses which non-clinical studies are necessary to demonstrate proof-of-principle and to define the pharmacological and toxicological effects predictive of the human response.

Special problems might be associated with the clinical development of human cell-based medicinal products. Guidance is therefore provided on the conduct of pharmacodynamic/pharmacokinetic studies, dose finding and clinical efficacy and safety studies. The guideline describes the special consideration that should be given to pharmacovigilance aspects and the risk management plan for these products.

1. INTRODUCTION (background)

Rapid development in the fields of biology, biotechnology and medicine has led to the development of new treatments and highly innovative medicinal products, including medicinal products containing viable cells. These new cell-based medicinal products have a high potential in the treatment of various diseases where there is a previously unmet medical need.

Human cell-based medicinal products are heterogeneous with regard to the origin and type of the cells and to the complexity of the product. Cells may be self-renewing stem cells, more committed progenitor cells or terminally differentiated cells exerting a specific defined physiological function. Cells may be of autologous or allogeneic origin. In addition, the cells may also be genetically modified. The cells may be used alone, associated with biomolecules or other chemical substances or combined with structural materials that alone might be classified as medical devices (combined advanced therapy medicinal products).

2. SCOPE

This multidisciplinary guideline will address development, manufacturing and quality control as well as non-clinical and clinical development of cell-based medicinal products (CBMP) including somatic cell therapy medicinal products as defined in Directive 2001/83/EC, Part IV, Annex I¹ and tissue engineered products as defined in Regulation 1394/2007/EC². This guideline is intended for products entering the Marketing Authorisation (MA) procedure. However, the principles laid down in the guideline should be considered by applicants entering into clinical trials.

Cell-based medicinal products discussed in this document have the following characteristics:

- They contain viable human cells³ of allogeneic or autologous origin undergoing a manufacturing process;

* Vigilance as described in Article 11 of Directive 2004/23/EC.

- They may be combined with non-cellular components;
- The cells may be genetically modified. The present document applies only to the cellular component of the cell based medicinal products containing genetically modified cells.

Although this document does not cover non-viable cells and cellular fragments originating from human cells, the underlying scientific principles may be applicable.

This guideline does not cover xenogeneic cell-based medicinal products.

3. LEGAL BASIS

This guideline should be read in conjunction with the introduction and general principles (4) and part 4 of the Annex I to Directive 2001/83/EC¹ as amended and the Regulation on Advanced Therapy Medicinal Products 1394/2007/EC².

Also, donation, procurement and testing of cells from human origin must comply with overarching Directive 2004/23/EC⁴ and technical directives drawn from it, Directives 2006/17/EC⁵ and 2006/86/EC⁶.

4. MAIN GUIDELINE TEXT

4.1 Risk analysis

The risk posed by the administration of a cell-based medicinal product is highly dependent on the origin of the cells, the manufacturing process, the non-cellular components and on the specific therapeutic use. The variety of cell-based medicinal products can lead to very different levels of risks for the patients, the medical personnel or the general population. Therefore the development plans and evaluation requirements need to be adjusted on a case by case basis according to a multifactorial risk based approach (see Annex I to Directive 2001/83/EC²).

At the beginning of the product development, an initial risk analysis may be performed based on existing knowledge of the type of product and its intended use. This should be updated by the applicant throughout the product life cycle as data are collected to further characterise the risk.

The comprehensive risk analysis should be used to justify the product development. It should also serve as a basis for the preparation of a risk management plan in accordance with the guideline on risk management systems for medicinal products for human use (EMA/CHMP/96268/2005)⁷. In particular, the results of the comprehensive risk analysis should be used:

- to identify risk factors associated with the quality and safety of the product;
- to determine the extent and focus of the data required during non-clinical and clinical development;
- when establishing the need for risk minimisation activities;
- when determining the post market risk management activities specified in the pharmacovigilance plan.

The following general risk criteria (non-exhaustive) can be used in the estimation of the overall risk of the product:

- origin (autologous-allogeneic);
- ability to proliferate and/or differentiate;
- ability to initiate an immune response (as target or effector);
- level of cell manipulation (in vitro/ex vivo expansion/activation/ differentiation /genetic manipulation/ cryo-conservation);
- mode of administration (e.g. ex vivo perfusion, local or systemic surgery);

- duration of exposure or culture (short to permanent) or life span of cell;
- combination product (cells and bioactive molecules or structural materials);
- availability of clinical data on or experience with similar products.

4.2 Quality and manufacturing aspects

This part of the guideline describes activities by manufacturers following procurement of the cells and tissues. The manufacture of cell-based medicinal products should be in compliance with the principles of good manufacturing practices, as set out in Directive 2003/94/EC⁸ and its Annex 2⁹.

The active substance of a cell-based medicinal product (CBMP) is composed of the engineered (manipulated) cells and/or tissues. Additional substances (e.g. scaffolds, matrices, devices, biomaterials, biomolecules and/or other components) when combined as an integral part with the manipulated cells are considered part of the active substance and are therefore considered as starting materials, even if not of biological origin.

CBMP often contain, or consist of cell samples of limited size and many are intended to be used in a patient-specific manner. This can raise specific issues pertaining to quality control testing designs for each product under examination. Since this document covers a variety of CBMPs, processes involved can vary from very simple to highly complex. For certain CBMPs, the starting material, the active substance and the finished product can be closely related or nearly identical. For such products, some requirements listed below could be inadequate and in those cases only relevant sections and items should be addressed.

4.2.1 Starting and raw materials

The manufacturing process of CBMP usually does not include terminal sterilisation, purification steps, viral removal and/or inactivation steps. Therefore, stringent sourcing requirements and acceptance criteria for all materials derived from human or animal origin should be adequately defined according to their intended use.

4.2.1.1. Cells

Donated cellular material from single or pooled donors, once processed (see 4.2.2.1) may be:

- A single primary cell isolate used directly for the CBMP;
- Primary cells cultured for a few passages before being used for the CBMP;
- Cells based on a well-defined cell bank system consisting of a master cell bank and a working cell bank.

An adequately controlled cell storage system should be established to allow proper maintenance and retrieval of cells without any alteration of their intended final characteristics. Storage conditions should be optimised to ensure cell viability, density, purity, sterility and function. Identity should be verified by relevant genotypic and/or phenotypic markers and the proportion of cells bearing these identity markers evaluated as an indicator of the intended cell population.

A. Cells of primary origin

The specific requirements for donation, procurement and testing laid down in Directive 2006/17/EC⁵ shall be met.

Procedures and standards employed for the selection of appropriate donors and the exclusion of high-risk or otherwise unsuitable candidate donors should be clearly delineated and justified. If it is necessary to pool cells from different donors, the risk analysis should address the possibility that pooling of allogeneic cell populations may increase the risk of undesired immunological responses in the recipient and compromise its therapeutic activity. In addition, pooling of cells may increase the

risk of disease transmission. Depending on the nature of the source of the cells and tissues, other risk factors, e.g. previous radiation exposure, should be also considered and addressed.

On receipt of the cells for use in a medicinal product, a specific microbiological screening programme should be in place, adapted to the type of cells, with validated assays capable of detecting human infectious agents with appropriate sensitivity and taking into consideration the medium components that might interfere with the assays (e.g. antibiotics). When cells originate from non-healthy tissues, the product specific acceptance criteria should be defined according to the intended use.

Quality parameters aimed at the definition of acceptance criteria for a given organ or tissues should be specified, taking into consideration general aspects such as shipment and storage conditions.

In the case of autologous donation, the testing regimen of the starting material should be justified, taking into account the autologous use.

Where allogeneic primary cells are collected and expanded for use in multiple patients, the cell lot should be appropriately characterised. The same characterisation programme shall be applied to each new cell lot.

B. Banking system for established cell lines

Where cell lines are used, an appropriately characterised Master Cell Bank (MCB) and Working Cell Bank (WCB) should be established, whenever possible. Cell banking and characterisation and testing of the established cell banks should comply with the ICH guideline Q5D¹⁰.

4.2.1.2. Other materials, reagents and excipients

Various materials are needed for collection, selection, culture or even genetic or phenotypic modification of cells, such as other cells, enzymes, antibodies, cytokines, sera and antibiotics. Exposure to such materials can also impact on the quality, safety and efficacy of the final therapeutic product. As a consequence, each substance used in the procedure should be clearly specified and evaluated as to its suitability for the intended use. The microbial purity and low endotoxin level of these materials should be ensured.

Materials, including cells that function as support for growth and adhesion e.g. feeder cells should be evaluated and/or validated as to their suitability for the intended use.

The quality of biologically active additives in culture media such as growth factors, cytokines and antibodies, should be documented with respect to identity, purity, sterility and biological activity and absence of adventitious agents. It is recommended to keep the use of such materials to a minimal and to avoid the use of reagents with sensitisation potential e.g. β -lactam antibiotics.

For viral safety aspects, the guidelines on viral safety^{11, 12} and Eudralex vol. 2B¹³ should be taken into consideration. The principles laid down in the general text of the European Pharmacopoeia on viral safety¹⁴ should be followed for every substance of animal and human origin that is used during the production.

Measures should be taken to reduce the risk of transmissible spongiform encephalopathy according to the relevant European legislation and guidelines¹⁵.

Where appropriate, the Note for Guidance on the “Production and quality control of medicinal products derived by recombinant DNA technology”¹⁶ and the Note for Guidance on the “Production and quality control of Monoclonal Antibodies”¹⁷ should be taken into account.

When the raw materials, reagents and/or excipients have a marketing authorisation or mentioned in a Pharmacopoeia, appropriate references may be given.

The following information must be added for materials of human or animal origin:

A. Human derived materials

Reagents of human origin (e.g. albumin, immunoglobulins) should be evaluated for their suitability in a manner identical to that employed for plasma-derived products as recommended in the CPMP Note for guidance on plasma-derived medicinal products¹⁸. The use of synthetic alternatives should be investigated. If serum is required in the culture media, the use of serum isolated from the same individual who donated the cells is preferred, where possible, to alternate allogeneic serum.

B. Animal derived material

Where cells or tissues of animal origin are used e.g. as supportive cells, the guidance given in “Points to consider on Xenogeneic Cell Therapy Medicinal Products”¹⁹ should be followed.

Animal derived reagents may harbour infectious agents and may increase undesirable immunological responses in the recipient. When applicable, the use of animal reagents should be avoided and replaced by non animal derived reagents of defined composition.

When bovine serum is used, the recommendations of the Note for Guidance on the “Use of Bovine Serum in the Manufacture of Human Biological Medicinal Product”²⁰ should be followed. The use of irradiated sera and/or alternative synthetic media is encouraged and should be considered.

For viral safety testing of materials of other animal species, the table of extraneous agents to be tested for in relation to the general and species-specific guidelines on production and control of mammalian veterinary vaccines²¹ and Note for Guidance on Production and Quality Control of Animal Immunoglobulins and Immunosera for Human use²² should be consulted.

C. Special considerations

Special recommendations for the starting materials of cell-based Gene Therapy Medicinal Products

When the cells in the active substance are genetically modified, the “Note for Guidance on the quality, preclinical and clinical aspects of gene transfer medicinal products”²³ should be followed, which gives details on the quality control, characterisation and preclinical testing of gene transfer vectors. Cell populations which are transformed should be assayed for appropriate and reproducible expression of the newly acquired characteristics. Special attention should be paid to the level and length of expression and quality of the gene product(s) produced by the cells. As far as applicable and practicable, the new characteristics of the cells should be quantified and controlled.

Special recommendations for matrix/device/scaffold components of combined products

Cell-based medicinal products may incorporate structural components which independently are medical devices or active implantable medical devices. Those devices should meet the essential requirements laid down in Directive 93/42/EEC²⁴ concerning medical devices and Directive 90/385/EEC²⁵ on the approximation of the laws of the Member States relating to active implantable medical devices, respectively, and this information shall be provided in the marketing authorization application. In the case where a Notified Body has evaluated the device part, the result of this assessment shall be included in the dossier. Cell-based medicinal products may also incorporate structural components which are not identical to, or used in the same way as in a medical device. All structural components should be appropriately characterised and evaluated for their suitability for the intended use (See sections on Characterisation and Development Pharmaceuticals).

Any matrices, fibers, beads, or other materials that are used in addition to or in combination with the cells should be described and their function underpinned by means of chemical, biological, physical (e.g. structure and degradation) and mechanical properties. Inclusion of additional bioactive molecules should also be described and their impact should be evaluated.

4.2.2 *Manufacturing process*

The manufacturing process of cell-based medicinal products should be carefully designed and validated to ensure product consistency. The requirements should be defined and justified.

A detailed description of the manufacture of the active substance and of the finished product should be provided. The type of manipulation(s) required for cell processing and the physiological function of the cells shall be described. A flow diagram of the entire process starting from biological fluid/tissue/organ or from cell banks should be prepared indicating critical steps and intermediate products (e.g. intermediate cell batches), as well as operating parameters, in-process controls and acceptance criteria. Manufacture of combined medicinal products consisting of cells and matrices/devices/scaffolds, require additional consideration regarding the cell-matrix/scaffold interactions and quality issues raised there from. Attention should be paid to biodegradable materials, which may possess the potential for environmental changes (e.g. raising pH) for the cells during the manufacture or after administration.

Information on procedures used to transport material during the manufacturing process of the product, including transportation and storage conditions and holding times, should be provided.

The manufacturing area should be physically separated from the procurement area. If different tissues and cellular products are processed and stored in the same manufacturing area there is an increased risk of cross contamination during each step of the procedure, e.g. via processing equipment or in storage containers such as liquid nitrogen tanks, and therefore, adequate control measures to prevent cross-contamination should be put into place.

Equipment and premises used for manufacturing of CBMP should be suitable and qualified for aseptic production. It is recommended that dedicated, product-specific or single-use equipment are used in the production, whenever possible.

1. Cell preparation procedures

All cell preparation procedures should be justified in terms of their intended purpose.

Inappropriate handling and improper processing of cells/tissues must be avoided as they can impair or destroy the integrity and/or function of the cells and thus result in therapeutic failure. Microbiological control is a pivotal aspect of process control and quality evaluation of all cell preparations. Monitoring of in vitro cell culturing at selected stages of the production should be performed where feasible. The culture should be examined for any microbial contamination in accordance with the culturing procedure and growth characteristics of the cells.

After appropriate controls have been performed / implemented, the biological fluid / tissue /organ can undergo one or more of the following steps:

Organ/tissue dissociation

The procedure to obtain the cells from the organ/tissue has to be described (with respect to the type of enzyme, media, etc.) and validated. Consideration should be given to the degree of disruption applied to the tissue in order to preserve the intended functional integrity of the cellular preparation and to minimize cell-derived impurities in the product (cell debris, cross contamination with other cell types).

Isolation of the cell population of interest

Any procedure used to isolate and / or purify the cell population of interest should be described. Its effectiveness should be addressed in relation to the intended use and the method(s) should be validated.

Cell culture

During in vitro cell culture, consideration should be given to ensure acceptable growth and manipulation of the isolated cells. The processing steps should be properly designed to preserve the integrity and control the function of the cells. The procedures for any manipulation should be documented in detail and closely monitored according to specific process controls. The duration of cell culture and maximum number of cell passages should be clearly specified and validated. The relevant genotypic and phenotypic characteristics of the primary cell cultures, of the established cell lines¹⁰ and the derived cell clones should be defined and their stability with respect to culture longevity determined. Consistency/repeatability of the cell culture process should be demonstrated and the culture conditions including the media and the duration should be optimised with respect to the intended clinical function of the cells.

Special consideration should be given to the growth potential of cells in response to growth factors since cell subpopulations may gain a growth advantage under defined in vitro culturing conditions.

Cell modification

Various treatments (physical, chemical or genetic) can be applied to cells. The method used to modify the cells should be fully described. In the case of genetic modification of cells, requirements set up in the Note for guidance on Quality, preclinical and clinical aspects of gene transfer medicinal products²³ should be followed.

Cells cultured in or on a matrix/device/scaffold

If the cells are grown directly inside or on a matrix/device/scaffold, the quality of the combined advanced therapy medicinal product relies predominantly on the properly controlled manufacturing process. For such products, the cell culture process has to be thoroughly validated and the effect of the device on the cell growth, function and integrity has to be taken into account. The effect that the cells may exert on the device (e.g. on rate of degradation) should also be considered (see also 4.2.6. Development Pharmaceuticals).

2. In-process controls

The manufacturing process needs to be controlled by several in-process controls at the level of critical steps or intermediate products. Intermediate cell products are products that can be isolated during the process; specifications of these products should be established in order to assure the reproducibility of the process and the consistency of the final product. Tests and acceptance criteria should be described. If storage occurs, it is necessary to validate the storage conditions (e.g. time, temperature).

3. Batch definition

The purpose of the batch definition is to ensure consistency and traceability. A clear definition of a production batch from cell sourcing to labelling of final container should be provided (i.e. size, number of cell passages/cell duplications, pooling strategies, batch numbering system). In the autologous setting, the manufactured product should be viewed as a batch.

4. Container and closure system

A description of the container closure system should be provided. Compatibility with the product should be demonstrated. It should be indicated if the container closure *per se* has a CE marking under the Medical Devices Directive 93/42/EEC²⁴. Information on the sterilisation procedures of the container and the closure should be provided.

The choice of packaging materials should be addressed as part of the development pharmaceuticals. Additional data may be required if packaging components are used in the transport and/or application procedure.

4.2.3 Characterisation

The characterisation of a CBMP should encompass all the components present in the finished product. Characterisation may prove particularly challenging for products containing cells together with matrices, scaffolds and innovative devices. Characterisation data are likely to be necessary for single components as well as for the combined final product. Characterisation data could encompass data obtained throughout the development and/or manufacturing process. It should be noted that in a combined product the characteristics of both the cellular and the non-cellular components may be altered by the process of integration.

An extensive characterisation of the cellular component should be established in terms of identity, purity, potency, viability and suitability for the intended use, unless justified.

The expected biological function of a CBMP encompasses complex interactions that may range from a biochemical, metabolic or immunological action to the structural replacement of damaged tissue or organ. Therefore, the requirements for a complete characterisation of the active substance in terms of biological function could be very taxing. Moreover the specific mechanism of action is often difficult to pinpoint to specific molecular entity but it is more dependent on the functionality of the cellular components acting in a “tissue-like” fashion as a whole. Therefore, when considering the extent of characterisation, the following issues should be taken into account: i) autologous cells vs. allogeneic cells, ii) extensively or minimally manipulated *in vitro*, iii) immunologically active or neutral, iv) proliferative capacity of the cells, v) cell-like or tissue-like organisation and dynamic interactions amongst cells and with the structural component, vi) intended use.

Non-cellular components should be characterised in the context of their required function in the finished product. This includes structural components designed to support the cellular components such as scaffolds or membranes which should be identified and characterised in chemical and physical terms such as porosity, density, microscopic structure and particular size according to the type of substances and intended use according to EN/ISO 10993-18²⁶ and EN/ISO 10993-19²⁷.

The characterisation should be designed to allow setting up the routine controls that will be applied for release of the active substance and finished product as well as those to be performed at several steps of the process to guarantee batch consistency.

If biologically active molecules (e.g. growth factors, cytokines etc.) are present as components of the cell-based products, these have to be described adequately and their interaction with the other components of the product and the surrounding tissues after administration should be characterised. This should involve an appropriate range of *in vitro* and where necessary *in vivo* methods.

1. Identity

Cellular Component

The identity of the cellular components, depending on the cell population and origin, should be characterised in terms of phenotypic and/or genotypic profiles.

When addressing the phenotype of the cells, relevant markers could be used, where justified. These markers may be based on gene expression, antigen presentation, biochemical activity, response to exogenous stimuli, capability to produce biologically active or otherwise measurable molecules, etc. For adherent cells, morphological analysis may be a useful tool in conjunction with other tests. Where applicable, a description of the procedures which could lead to a modification of the characteristic of the product, including adhesion, absorption, degradation, presentation of components of the culture media, should be provided.

For cellular components of allogeneic origin, identity should include histocompatibility markers, where applicable, and identification of genetic polymorphisms with specific reference to the intended use.

Non-cellular Components of the active substance

All non-cellular components should be appropriately characterised as such and identity parameters established.

Should the finished product contain a distinct active substance in addition to the cellular component, then that active substance should be characterised with respect to identity in accordance to relevant CHMP guidelines, depending on the nature of the active substance, whether it be of chemical or biological origin.

Structural components designed to support the cellular components such as scaffolds or membranes should be identified and characterised with respect to their composition and structural characteristics.

Combination Products

In a combination product the active substance may be formed by the integration of cellular and non-cellular components to form a single entity. In such a case the identity of both the cellular and the non-cellular components may be altered by the process of combination. Consequently a distinctive way to define identity should be established for the components in the combination, unless justified.

2. Cell purity

The cellular population of interest could contain other cells that are of different lineages and/or differentiation stage or that may be unrelated to the intended population.

Where a specific cell type is required for the indication, the unwanted cells should be defined and their amount in the final product should be controlled by appropriate specifications, i.e. acceptance criteria for the amounts of contaminating cells should be set.

In cases, where the desired biological activity and efficacy of the product requires a complex mixture of cells, the cell mixture needs to be characterised and its composition controlled by appropriate in-process controls and release testing.

Irrespective of the cell type, the cell population can be contaminated with non-viable cells. Since cell viability is an important parameter for product integrity and directly correlated to the biologic activity, the ratio between non-viable and viable cells should be determined and specifications should be set.

3. Impurities

Product or process-related

During the production of a CBMP, variable amounts of impurities, product- and process-related, may be introduced into the final product. Any reagents known to be harmful in humans should be analysed in the final product (or in individual components if otherwise not possible) and acceptance criteria should be set. The specification limits should be justified by levels detected in batches used for toxicological and/or clinical studies.

Any material capable to introduce degradation products into the product during the production, e.g. biodegradable materials, should be thoroughly characterised in this respect and the impact of the degradation products to the cell component(s) should be addressed.

If genetically modified cells are used in the product, any additional proteins expressed from the vector, e.g. antibiotic resistance factors, selection markers, should be analysed and their presence in the product should be justified.

Adventitious agents

A critical aspect is to establish that CBMP are free from adventitious microbial agents (viruses, mycoplasma, bacteria, fungi). The contamination could originate from the starting or raw materials (see above), or adventitiously introduced during the manufacturing process. A risk assessment should be performed to evaluate the possibility of reactivation of cryptic (integrated, quiescent) forms of adventitious agents. A thorough testing for the absence of bacteria, fungi and mycoplasma shall be performed at the level of finished product. These tests should be performed with the current methodologies described in the European pharmacopoeia for cell based products²⁸. In cases where the short shelf life of the CBMP is prohibitive for the testing of absence of bacteria under the Eur Ph. requirements, alternative validated testing methods may be acceptable, if justified.

4. Potency

It is strongly recommended that the development of a suitable potency assay be started as soon as possible. Preferably, a suitable potency assay should already be in place when material for the first clinical trial is produced and it should be validated prior to pivotal clinical trials unless otherwise justified. Lot release and shelf life specifications for potency should be determined and amended during product development, if appropriate.

According to the ICH guideline 6QB²⁹, potency is the quantitative measure of biological activity based on the attribute of the product, which is linked to the relevant biological properties. The assay demonstrating the biological activity should be based on the intended biological effect which should ideally be related to the clinical response.

Basically, two types of potency assays can be envisioned: 1) *in vitro* assays using cell systems and 2) *in vivo* assays using animal models. Major cellular functions as viability, self renewal, death and differentiation are pivotal to the quality, function and sustainability of the CBMP and may need to be monitored during production and at release using surrogate markers and appropriate technology (e.g. gene expression profiles by microarrays, flow cytometric immunofluorescent analysis, cell cloning, PCR and many others). *In vivo* assays for potency may also be useful especially when experimental animal models are available.

Markers for purity and markers for potency should not be mixed in the same assay. .

Reference is made to the Guideline on “Potency testing of cell-based immunotherapy medicinal products for the treatment of cancer”³⁰. Although this guideline focuses on cell-based immunotherapy medicinal products, the principles, including on reference preparations, apply for all CBMP. As described in the referred guideline, a combination of multiple methods may be needed to adequately define the potency of these products during the development. Certain assays may be needed to control process changes, whereas others are more suitable for release testing.

Tissue repair and regeneration

An *in vivo* test can either be performed in an animal model mimicking the intended clinical tissue repair/ regeneration or can otherwise be based on the mode of action (e.g. an ectopic model). An *in vitro* assay can be based on the expression of markers that have been demonstrated to be directly or indirectly (surrogate markers) correlated to the intended biological activity, such as cell surface markers, activation markers, expression pattern of specific genes. Also a physiological response under defined conditions such as differentiation in specific cell types and/or secretion of tissue specific proteins (e.g. extracellular matrix components) can be used as a basic principle for a potency test. The manufacturers should, however, ensure that the method of characterisation is relevant for the intended biological effect *in vivo*.

The potency assay should be performed by using a specified number of cells and, when possible, quantified against a qualified reference preparation. The potency should be defined as the required time to obtain a predefined effect (e.g. restoration of function or repair of anatomical structure) or the potency is calculated from the measured effect in a defined time period.

Metabolic or pharmacological activity

Cells contained in a CBMP can be chemically treated or genetically modified *in vitro* to express certain desirable proteins like growth factors, cell surface antigens or other molecules in order to sustain the biological response as long as needed in the new microenvironment. Therefore, the potency assays to be developed should be able to assess the activity-related aspects of the active substance that may be composed not entirely of intact viable cells but also of other components.

If the intended biological function of the CBMP is mainly based on the capacity of cells to secrete specific molecule(s) e.g. to repair a metabolic disorder, to promote growth, to release a metabolite, then its potency assay will be based on the detection of the active molecule(s) produced and the biological activity expected. This can be carried out by conventional reliable qualitative and quantitative analytical methods (protein analysis, nucleic acid identification, HPLC chromatography etc.). The same molecule can be also assessed for function in animal model systems assuming that the active substance is released from the cell-based medicinal product into biological fluids (plasma, CSF, urine or interstitial fluid).

Immunotherapy

Potency assays of cell-based medicinal products intended for immunotherapeutic use will be based on complex immune mechanisms which may be complicated by multi-antigen formulations and inherent variability of the starting material. Special guidance for cell-based immunotherapy medicinal products is provided in Guideline on “Potency testing of cell-based immunotherapy medicinal products for the treatment of cancer”³⁰.

5. Tumourigenicity

The tumourigenicity of CBMP differs from the classical pharmaceuticals as the transformation can also happen in the cellular component of the product (e.g. chromosomal instability) and not only in the treated individual. If the risk of cellular transformation and subsequent potential for tumourigenicity can be foreseen, the cellular components should be evaluated for their tumourigenic potential by analysing e.g. their proliferative capacity, dependence on the exogenous stimuli, response to apoptosis stimuli and genomic modification. Testing of chromosomal integrity and tumourigenicity of cells derived from a cell culture / cell banking system will be required. Reference is made to the ICH Q5D¹⁰ and to the Ph. Eur. Monograph on cell substrates for the production of vaccines for human use³¹.

4.2.4 Quality control

For proper quality control, the active substance and/or the final product should be subjected to release testing, whenever possible. If justified, it would be acceptable to have reduced testing at one level provided an exhaustive control is performed at the other. All release testing should be performed using methods validated at the latest at the time of submission of an application.

1. Release criteria

The release specifications of the active substance and finished product should be selected on the basis of parameters defined during the characterisation studies. Selection of tests is product-specific and has to be defined by the manufacturer.

Specifications for release testing should include identity, purity, potency, impurities, sterility, potency, cell viability and total cell number, unless otherwise justified. If the structure is an essential characteristic of the product, the structural characteristics of the active substance or finished product shall be defined and justified. In case the primary function of the CBMP is the excretion of specific proteins, specifications regarding these excreted proteins should be set.

If certain release tests cannot be performed on the active substance or finished product, but only on key intermediates and/or as in-process tests, this needs to be justified. In these cases an adequate

quality control has to rise from the manufacturing process, supported by the results of the clinical studies. These exceptions may include the following:

- Some release tests might not be feasible on the combined components of the active substance/ finished product for technical reasons.
- A complete release testing cannot be finalised before the product is administered to the recipient due to time restrictions (e.g. in case of autologous products, which are administered immediately after completion of the production and initial testing). However, a critical set of essential tests that can be performed in the limited time prior to clinical use must be defined and justified. Whenever feasible, retention samples should be stored for future analysis.
- The amount of available product is limited to the clinically necessary dose (e.g. due to very limited cell numbers at collection or low proliferation rates). The release of the product should be justified by the validation of the cell manipulation process and the in-process controls.

2. Stability testing

A shelf life for the cells under specified storage conditions shall be determined for the following materials: i) all intermediates subject to storage if applicable, ii) components of the combined CBMP, iii) the active substance, iv) the finished product. Furthermore, a valid in-use shelf life (after opening from the transport container) should be assigned to the CBMP. Also, all storage conditions including temperature range should be defined. Transportation and storage conditions should be supported by experimental data with regard to the maintenance of cell integrity and product stability during the defined period of validity. If relevant, appropriate methods for freezing and thawing should be documented.

Due to the complex nature of the active substance of a CBMP, requirements for stability should be defined on a case-by-case basis. Whenever possible, stability should be assessed for both the cellular as well as the non-cellular component prior to combination and together as a finished product in the final packaging.

3. Special quality requirements for cell-based medicinal products containing genetically modified cells

If cells have been genetically modified, quality control must be performed in compliance with guidance available on gene transfer medicinal products²³. This information is in addition to control of the cells according to the guidance presented elsewhere.

4. Special quality requirements for combination products

Specifications for structural components of the product shall be defined. Impurities and degradation product that originate from the structural component (matrix, scaffold, device) shall be described and specifications for the relevant impurities should be set. Testing of the structural/mechanical properties and biological activity with reference to the anticipated conditions for use and potential for degradation may be difficult to conduct as part of release testing. Thus, it is anticipated that these parameters could be explored through proper testing of raw materials and characterisation studies of the final product. In extremely limiting conditions (e.g. for autologous products with small cell numbers), the analysis of structural/functional characteristics of a combination product may necessitate the development of a model product composed of same non-cellular components combined with cell component(s) of equal characteristics but with proven availability.

4.2.5 Validation of the manufacturing process

The entire manufacturing process, including cell harvesting, cell manipulation processes, maximum number of cell passages, combination with other components of the product, filling, packaging, transport, storage etc., should be validated. Validation of the production process of a combined

product should encompass all steps from separate components up to the final combination to ensure consistent production.

It should be demonstrated that each step of the manufacturing process of the active substance, supportive components and final product is well controlled. The selection and acceptance criteria of the operational parameters and the in-process controls should be justified. Putative variability, related to starting materials and biological processes, should be taken into account in the validation. Furthermore, the critical points of the manufacturing process should be defined and validated, especially the aseptic processing.

Any preservation steps, holding periods and/or transportations of the active substance, final product, supportive structures or intermediate products during the manufacturing process should be validated.

In case of limited sample sizes (e.g. autologous preparations for one single administration), it is recommended that a more extensive validation is performed with cell preparations of comparable characteristics but available in sufficient amounts for validation purposes. It is recommended that validation of such a manufacturing process is performed depending on the product characteristics, for adventitious agents, identity, potency, viability, purity/impurities and other product specific parameters.

4.2.6 *Development Pharmaceuticals*

The general principles set out in Note for Guidance on Development Pharmaceuticals for Biotechnological/Biological Products³² can be applied to human CBMP. The potential complexity of composition and the dynamic nature of a product containing living cells will result in very specialised pharmaceutical and biopharmaceutical requirements for each development programme from the individual cell components into the final product.

1. Cellular Components

The development programme should address the choices of materials and processes to be used in production. This should be addressed from the point of view of the biological/therapeutic function, the maintenance and the protection of the cell population.

Integrity of the cellular component is most critical for the CBMP and must be assessed by the ability of cells to survive, and maintain the genotype or phenotype needed for the intended functions. However, detection of possible changes in cellular nature that may influence the intended function, can be feasible by analysis of cellular surface antigens, proteomics and functional genomics analysis (e.g. microassay for gene expression profile, flow cytometry etc.). Cell viability can be easily assessed in culture by employing widely applied assays. For combined products, where structural components are an integral part of the active substance, such assays may be more difficult to apply. Alternative approaches could be sought such as combination of other suitable assays (e.g. detection of pH and O₂ / CO₂), where needed.

The ability of cells to continue to produce or express products should be evaluated as part of the stability programme. Such stability studies should be carried out as long as the defined period of validity requires.

2. Non-Cellular Components

A CBMP may contain non-cellular components, such as biomaterials, bioactive molecules, proteins or chemical entities. These may supply structural support, suitable environment for growth, biological signalling or other functions. They may also be used during the *ex vivo* manipulation process.

Matrices, scaffolds, devices, biomaterials or biomolecules which are not an integral part of the active substance, are considered as excipients of the finished product. For excipient(s) used for the first time in combination with cells and/or tissues, the requirements for novel excipients, as laid down in part I,

Annex I of Directive 2001/83/EC¹, do apply. Conventional excipients should also be characterised with respect to their combination with cells.

Information on the choice of excipients, their properties, their characteristics and the design and testing of a final scaffold/matrix should be provided in the dossier as part of the development pharmaceuticals.

Should the finished product contain components that act to modify the delivery or ensure the local retention of cells after administration, the scientific rationale should be provided and supported by adequate development data. The evaluation of individual non-cellular components is required although aspects of this evaluation may be incorporated into studies designed to assess the product as a whole. Where the safety of a non-cellular component has previously been established for other applications, for example in support of the approval of a particular material for a medical device or medicinal product application, elements of that evaluation may be applicable to an evaluation of its safety and suitability when used in a cell-based medicinal product, if justified.

The relevance of the structural and functional characteristics of the non-cellular components in a combination product should be discussed. Interaction of the cellular component and any additional non-cellular components with the device should be evaluated and the development and characteristics of the combined product as a whole should be presented.

Tissue differentiation and functionality are highly dependent on the local environment and thus on the choice of biomaterials and cell signalling biomolecules (e.g. growth factors). Therefore, studies should be carried out to verify critical aspects of the character and performance of biomaterials and other non-cellular components used in the CBMP, for example biocompatibility and mechanical strength.

In particular, to confirm that the properties of a biomaterial permit the growth and proper function of the tissue/cells with which it is in contact and support the overall performance of the product, assurance should be provided in relation to the following:

- absence of components or leachables that might be toxic to cell growth and/or to the intended performance;
- characterisation of features (e.g. topography, surface chemistry, strength) critical to structural support, optimisation of viability and cellular growth or other functional characteristics;
- biocompatibility of the structural material with the cells or tissues to confirm that the system maintains the desired cell differentiation, functionality and genotype during production and until use;
- release kinetics and/or rate of degradation of any bioactive molecules, to verify that they are appropriate for the achievement of the intended effect.

To establish biocompatibility, it is necessary to specify the nature of biological responses that a biomaterial is required to elicit from the host tissue or cell-based components, and to provide evidence that the desired tissue response is achieved in a relevant model.

The stability of the non-cellular components should be assessed in the presence and absence of cellular components in order to determine whether the non-cellular component undergoes degradation, or physico-chemical alterations (e.g. aggregation, oxidation) that may impact on the quality of the product by affecting cellular behaviour and survival. The effect of the cellular component or of the surrounding tissues on the degradation (rate and, if appropriate, products) or stability of the structural component should be assessed, considering also the effect of the non-cellular components throughout the expected lifetime of the product. The general principles that are applied to the biological evaluation of medical devices can also be applied to the evaluation of biomaterials intended for use in CBMP. Such an evaluation involves a programme of characterisation, testing and review of existing data to assess the potential for an adverse biological reaction to occur as a result of exposure to the biomaterial. These principles are set out in international standard ISO 10993 Part 1³³. Other parts of the ISO 10993 series of standards specify methods that may be relevant to the assessment of material characteristics, biological safety and degradation of biomaterials used in cell-based medicinal

products. Additional studies (e.g. cell adhesion studies, growth studies) may be necessary to demonstrate aspects of biocompatibility specific to cell-based applications.

3. Final Product

Once the “formulation”, i.e. the delivery system of a combined product has been established, the parameters for determining the role of constituents and appropriateness of composition should be presented within a justification of the composition of the product.

The key parameters for performance testing of the completed product should be justified in relation to the development data and the final quality requirements. It may be appropriate that *in vitro* and *in vivo* testing of the formulation/delivery system/combined product during development are included.

4.2.7 Traceability

A system allowing complete traceability of the patient as well as the product and its starting materials is essential to monitor the safety and efficacy of cell-based medicinal products. The establishment and maintenance of that system should be done in a way to ensure coherence and compatibility with traceability and vigilance requirements laid down in Directive 2004/23/EC⁴ and in Directives 2006/17/EC⁵ and 2006/86/EC⁶ and Art. 15 of the Regulation (EC) No 1394/2007².

Article 15 of the Regulation (EC) No 1394/2007² defines a two tiered system connecting the required traceability from cell donation and procurement (Dir 2004/23/EC) to the manufacturer and user (hospital or practice). This means that anonymity can be guaranteed. At the tissue establishment there has to be a link between the donor and the donation. At the manufacturing side, there has to be a link between donation and product and at the hospital/practice side there has to be a link between the product and the recipient. The systems should allow full traceability from the donor to the recipient through anonymous coding systems. Manufacturers should establish their coding systems in a rational way, building from the coding system of the tissue establishment, and designing it to facilitate the tracing of the donation to the product and to the patient. Bar coding and peeling labelling systems could be suitable tools for the purpose of patient management.

4.2.8 Comparability

Development of a cell-based medicinal product may encompass changes in the manufacturing process that might have an impact on the final product. Given the complex and dynamic nature of CBMP it is particularly important that all stages of development are fully evaluated and tracked within the dossier. This is especially significant once clinical studies have commenced. Data on the behaviour and characteristics of developmental prototypes should be retained as it could provide background information relevant to the evaluation of the final product. During the pivotal clinical studies changes should not be introduced to the manufacturing process and the final product.

Materials used in the clinical studies should be sufficiently characterised in order to allow the demonstration of consistency in the production. The manufacturers should consider the critical parameters drawn from the characterisation of their product to establish the analytical tools necessary for the required comparability studies throughout development. Comparability studies with the product resulting from those changes should be performed in relation to clinical trial batches that were used. Appropriate guidance can be found in ICH Q5E Comparability of Biotechnological/Biological Products³⁴ and related guidance documents.

Whenever comparability at the analytical and/or non-clinical level cannot be established, it must be demonstrated by clinical data.

4.3 **Non-clinical development**

The scrutiny applied during non-clinical testing should take into account the nature of the Cell-based medicinal product (CBMP) and be proportional to the risk expected to be associated with clinical use.

The variability of cell-based medicinal products should be reflected in the non-clinical studies. Conventional requirements as detailed in Module 4 for pharmacological and toxicological testing of medicinal products may not always be appropriate. Any deviation from these requirements shall be justified. If cells in a CBMP have been genetically modified, non-clinical development must be performed in compliance with guidance available on gene transfer medicinal products²³.

The objectives of the non-clinical studies are to demonstrate proof-of-principle, define the pharmacological and toxicological effects predictive of the human response, not only prior to initiation of clinical trials, but also throughout clinical development. The goals of these studies include the following: to provide information to select safe doses for clinical trials, to provide information to support the route of administration and the application schedule, to provide information to support the duration of exposure and the duration of the follow-up time to detect adverse reactions, to identify target organs for toxicity and parameters to monitor in patients receiving these therapies.

The non-clinical studies should be performed in relevant animal models. If relevant animal models cannot be developed, *in vitro* studies may replace animal studies. The rationale underpinning the non-clinical development and the criteria used to choose a specific animal model must be justified. Expression level of biologically active molecules, the route of administration and the dosages tested should reflect the intended clinical use in humans

The recommendations of the ICH S6 Guideline³⁵ should be considered. The number of animals, their genders, the frequency and duration of monitoring should be appropriate to detect possible adverse effects.

The safety and suitability of all structural components for their intended function must be demonstrated, taking into account their physical, mechanical, chemical and biological properties. (See section 4.2.6 Development pharmaceuticals).

4.3.1. *Pharmacology*

Primary pharmacodynamics

Non-clinical studies should be adequate to demonstrate the proof of principle of the CBMP. The principal effects should be identified in non-clinical studies in a suitable model *in vitro* or *in vivo*.

Reasonably justified markers of biological activity should be used to adequately identify the pharmacodynamic action of the CBMP in the host.

If the intended use of the CBMP is, for example, to restore the function of deficient cells/tissue (tissue regeneration), functional tests should be implemented to demonstrate that function is restored. If the intended use is, for example, adoptive immunotherapy in cancer patients, the biological effect should be supported by data describing the immunologic action of the CBMP.

The chosen animal model may include immunocompromised, knockout or transgenic animals. Homologous models may be advantageous, since the *in vivo* behaviour of the applied cells or tissue in heterologous models could be altered due to species-specific mismatches. Homologous models should be considered for the study of stem cell differentiation. *In vitro* studies, addressing cell and tissue morphology, proliferation, phenotype, heterogeneity and the level of differentiation may be part of the primary pharmacodynamic analyses.

If possible, studies should be conducted in order to determine the minimal or optimal effective amount of CBMP that is needed to achieve the desired effect.

Secondary pharmacology

Potential undesirable physiological effects of human CBMP including their bioactive products should be investigated in an appropriate animal model. Cells may migrate from their intended location and, after a systemic administration, may home to other organs beside the intended location. Also, somatic cells may secrete additional biologically active molecules besides the protein of interest. Also, the protein(s) of interest can have additional targets beside the desired one.

Safety pharmacology

Safety pharmacology should be considered on a case-by-case basis depending on the characteristics of the CBMP. Cells may secrete pharmacologically active substances resulting in CNS, cardiac, respiratory, renal or gastrointestinal dysfunctions. Alternatively, cells by themselves could be envisaged to induce such consequences for example stem cells or muscle cells transplanted to infarcted regions of the heart.

For further guidance see ICH S7A Guideline³⁶ when applicable.

Kinetics, migration and persistence

Conventional ADME studies are usually not relevant for human CBMP. However, studies should be carried out to demonstrate tissue distribution, viability, trafficking, growth, phenotype and any alteration of phenotype due to factors in the new environment.

Cells may migrate within the host, thus presenting clinical concerns regarding adverse reactions deriving from displaced, possibly differentiating cells. This should be evaluated in animals using appropriate methods for specific identification of the cells.

Regarding biodistribution, the use of small animals allows meticulous cell detection, which will be practically more difficult in larger animals.

For human CBMP producing systemically active biomolecules, the distribution, duration and amount of expression of these molecules and the survival and the functional stability of the cells at target sites should be studied.

Interactions

The interaction of the applied cells or surrounding tissue with the non-cellular structural components and other bioactive molecules as well as the integration of the CBMP with the surrounding tissue should be monitored.

4.3.2. Toxicology

The need for toxicological studies depends on the product. However, as conventional study designs may not be appropriate, the scientific justification for the models used, or the omission of studies, shall be provided.

Toxicity may evolve, for example, due to unknown cellular alterations developing during the manufacturing process such as altered excretion patterns and *in vivo* behaviour due to differentiation of the cells. Other potential factors that may induce toxicity include the allogeneic use of the product, the presence of components that are used in the manufacturing process or are part of a structural component, or proliferation of the applied cells in an unwanted quantity or in an unwanted location.

Conventional toxicology studies might nevertheless be required, for example for complex regimens where CBMP are combined with other medicinal products or treatments such as adjuvants/cytokines or irradiation, respectively. The need for drug interaction studies is dependent on the intended use and the type of the cell-based product and should be discussed.

The induction of an immune response against the cells themselves and/or towards cell-derived pharmacologically active substances might modulate the efficacy of the CBMP. Therefore, the possible immunogenicity of a CBMP should be considered. For guidance on immunogenicity of excreted substances see ICH S6 Guideline³⁵.

Auto-immunity should be considered when cells are used for immunotherapy purposes, e.g. cancer immunotherapeutic products.

Single and repeated dose toxicity studies

Toxicity studies should be performed in relevant animal models. If the human cells are not immediately rejected, the studies may be combined with safety pharmacology, local tolerance, or proof of concept and efficacy studies. Sufficiently characterized analogous animal-derived cells may be used for some allogeneic CBMP when not immediately rejected.

The duration of observations in such studies might be much longer than in standard single dose studies, since the cells are supposed to function for long times, or induce long term effects, which should be reflected in the design of these studies. The route and dosing regimen should reflect the intended clinical use. Repeated dose toxicity studies are only relevant if the clinical use includes multiple dosings.

Local tolerance studies

Local tolerance studies may be required in an appropriate species. Most often, local tolerance, tissue compatibility and tolerance to excreted substances can be evaluated in single or repeated dose toxicity studies.

Other toxicity studies

The risk of inducing tumourigenesis due to neoplastic transformation of host cells and cells from the CBMP should be considered, as appropriate, on a case-by-case basis. Conventional carcinogenicity studies may not be feasible. Tumourigenesis studies should preferably be performed with cells that are at the limit of routine cell culturing or even beyond that limit. Tissues found to contain applied cells or expressed products during the biodistribution studies should also be analysed with special emphasis during tumourigenicity studies.

Genotoxicity studies are not considered necessary for human CBMP, unless the nature of any expressed product indicates an interaction directly with DNA or other chromosomal material.

The need for reproductive studies is dependent on the CBMP and should be considered on a case-by-case basis.

4.4 Clinical development

4.4.1 General aspects

In general, when a Cell-based medicinal product (CBMP) enters the clinical development phase, the same requirements as for other medicinal products apply. The clinical development plan should include pharmacodynamic studies, pharmacokinetic studies, mechanism of action studies, dose finding studies and randomised clinical trials in accordance to the Directive 2001/20/EC³⁷ and to the existing general guidances and specific guidances for the condition evaluated.

Due to specific biologic characteristics of CBMP, alternative approaches to Phase I to Phase III clinical trials might be required and acceptable for clinical development, if justified. The relevant non-clinical studies, previous clinical experience of the treated pathology, and initial clinical studies could be applied for demonstration of the “proof of principle” and the choice of clinically meaningful endpoints for safety and efficacy evaluation. European scientific advice is strongly recommended in case of uncertainties in rationale of CBMP development at the earliest phase.

CBMP might require administration through specific surgical procedures, method of administration or the presence of concomitant treatments to obtain the intended therapeutic effect. The biological effects of CBMP are highly dependent on the *in vivo* environment, and may be influenced by the replacement process or the immune reaction either from the patient or from the cell-based product. These requirements coming from the clinical development should be taken into account for the final use of these products. Their standardisation and optimisation should be an integral part of the clinical development studies. The therapeutic procedure as a whole, including the method of administration and required concomitant medication, such as immunosuppressive regimens need to be investigated and described in the product information, notably in the Summary of Product Characteristics (SPC).

4.4.2 Pharmacodynamics

Even if the mechanism of action is not understood in detail, the main effects of the CBMP should be known. When the purpose of the CBMP is to correct the function of deficient or destroyed cell/tissue, then functional tests should be implemented. If the intended use of the CBMP is to restore/replace cell/tissues, with an expected lifelong functionality, structural/histological assays may be potential pharmacodynamic markers. Suitable pharmacodynamic markers, such as defined by microscopic, histological, imaging techniques or enzymatic activities, could be used.

When CBMP includes a non cellular component, the combination should be assessed clinically for compatibility, degradation rate and functionality.

4.4.3 Pharmacokinetics

Conventional ADME studies are usually not relevant for human CBMP. Study requirements, possible methodologies and their feasibility shall be discussed, attention being paid to monitoring of viability, proliferation/differentiation, body distribution / migration and functionality during the intended viability of the products.

If multiple (repeated) administrations of the CBMP are considered, the schedule should be discussed in view of the expected *in vivo* life span of the CBMP.

4.4.4 Dose finding studies

The selection of the dose should be based on the findings obtained in the quality and the non-clinical development of the product and it should be linked with the potency of product. Even though the dosage for cell-based product might be determined by individual characteristics of the intended

patients (i.e. cell mass density per body weight/ volume of missing tissue/ missing surface), the dose to be tested in the confirmatory trial should be supported by the evidence provided by the phase I/II studies.

Phase I/II studies should be designed to identify a Minimal Effective Dose, defined as the lowest dose to obtain the intended effect or an Optimal Effective Dose Range, defined as the largest dose range required to obtain the intended effect based on the clinical results for efficacy and tolerability. If possible, also the Safe Maximal Dose, defined as the maximal dose which could be administered on the basis of clinical safety studies without acceptable adverse effects, should be investigated.

4.4.5 *Clinical Efficacy*

Clinical efficacy studies should be adequate to demonstrate efficacy in the target patient population using clinically meaningful endpoints, to demonstrate an appropriate dose-schedule that results in the optimal therapeutic effect, to evaluate the duration of therapeutic effect of the administered product and to allow a benefit – risk assessment taking into account the existing therapeutic alternatives for the target population. Confirmatory studies should be, as stated before, in accordance to the existing general guidelines and specific guidelines for the condition evaluated.

Deviations from these will need a justification. For example, the fact that the nature and the mechanism of action of the CBMP may be entirely novel does not mean necessarily that the therapeutic benefit should be measured by different endpoints from those recommended in the current disease-specific guidelines (e.g. medicines vs. cell implants for Parkinson's disease).

For new therapeutic applications of CBMP where limited guidance exists, consultation of regulatory authorities on the clinical development plan, including the confirmatory studies, is highly recommended.

The use of previously validated or generally accepted surrogate endpoints is possible provided that a correlation-between clinical meaningful endpoints and efficacy can be established. Sometimes, the desired clinical endpoint, such as prevention of arthrosis, can be observed only after a long follow up. In such cases, the marketing authorisation can be based on surrogate markers. If the efficacy is dependent on the long-term persistence of the product, a long-term follow up plan of the patients should be provided. Thus, the use of novel meaningful endpoints, clinical or other, is acceptable if justified.

4.4.6 *Clinical Safety*

The safety database should be able to detect common adverse events. The size of the database might be decided also in the light of previous clinical experience with similar products.

The risk of the therapeutic procedure as a whole, e.g. the required surgical procedures to administer the CBMP or the use of immunosuppressive therapy, shall be evaluated and used to justify the clinical studies and the choice of the target patient population

All safety issues arising from the preclinical development should be addressed, especially in the absence of an animal model of the treated disease or in the presence of physiologic differences limiting the predictive value of homologous animal model.

Particular attention should be paid to those biological processes including immune response, infections, malignant transformation and concomitant treatment during development and post-marketing phase of cell-based medicinal products.

For products with expected long term viability, patient follow-up is required in order to confirm long term efficacy and safety issue related to the product.

Clinical safety studies on repeated administrations should be performed as required by the risk analysis. The definition of Maximal Safe Dose should take into account also the possibility of repeated administration.

4.4.7 Pharmacovigilance and Risk Management Plan

The routine pharmacovigilance and traceability of the product should be described in the EU Risk Management Plan (RMP) as described in Guideline on risk management systems for medicinal products for human use⁷. CBMP may need special long-term studies to monitor specific safety issues, including loss of efficacy.

The long-term safety issues, such as infections, immunogenicity/immunosuppression and malignant transformation as well as the *in vivo* durability of the associated medical device/biomaterial component should be addressed in the RMP. Special pharmacoepidemiological studies may be needed. The specific requirements are linked to the biologic characteristics of the cell-based product. Traceability in the donor-product-recipient axis, or of the product-recipient for autologous products, is required in all circumstances as described in Directive 2004/23/EC⁴ and in Regulation (EC) No. 2007/1394/EC on Advanced Therapy Medicinal Products².

REFERENCES (scientific and / or legal)

¹ Directive 2001/83/EC of the European Parliament and of the Council of 6 November 2001 on the Community code relating to medicinal products for human use.

² Regulation (EC) No 1394/2007 of the European Parliament and of the Council on advanced therapy medicinal products and amending Directive 2001/83/EC and Regulation (EC) No 726/2004.

³ Ph. Eur. General Chapter 2.7.29: *Nucleated cell count and viability*. (01/2008:20729)

⁴ Directive 2004/23/EC of the European Parliament and of the Council of 31 March 2004 on setting standards of quality and safety for the donation, procurement, testing, processing, preservation, storage and distribution of human tissues and cells.

⁵ Commission Directive 2006/17/EC of 8 February 2006 implementing Directive 2004/23/EC of the European Parliament and of the Council as regards certain technical requirements for the donation, procurement and testing of human tissues and cells.

⁶ Commission Directive 2006/86/EC of 24 October 2006 implementing Directive 2004/23/EC of the European Parliament and of the Council as regards traceability requirements, notification of serious adverse reactions and events and certain technical requirements for the coding, processing, preservation, storage and distribution of human tissues and cells.

⁷ EMEA/CHMP Guideline on risk management systems for medicinal products for human use (EMA/CHMP/96268/2005)

⁸ Directive 2003/94/EC laying down the principles and guidelines of good manufacturing practice in respect of medicinal products for human use and investigational medicinal products for human use.

⁹ Annex 2 of Directive 2003/94/EC: Manufacture of Biological Medicinal Products for Human Use.

¹⁰ ICH Q5D, Derivation and Characterisation of Cell Substrates Used for Production of Biotechnological/Biological Products (CPMP/ICH/294/95)

¹¹ ICH Q5A Guideline on Quality of Biotechnological Products: Viral Safety Evaluation of Biotechnology Product Derived From Cell Lines in of human or animal origin (CPMP/ICH/295/95)

- ¹² EMEA/CPMP Note for Guidance on Virus Validation Studies: The Design, Contribution and Interpretation of Studies validating the Inactivation and Removal of Viruses (CPMP/BWP/268/95)
- ¹³ Eudralex Vol. 2 B, Notice To Applicant, part II-V : virological documentation
- ¹⁴ Ph. Eur. General Text 5.1.7: Viral safety (01/2008:50107)
- ¹⁵ EMEA/CPMP/CVMP Note for guidance on minimizing the risk of transmitting animal spongiform encephalopathy agents via human and veterinary medicinal products (EMA/410/01 rev.2)
- ¹⁶ Guideline on Production and Quality Control of Medicinal Products Derived by Recombinant DNA Technology. (3AB1A)
- ¹⁷ EMEA/CHMP Note for Guidance on Production and quality control of Monoclonal Antibodies (CHMP/BWP/157653/07)
- ¹⁸ EMEA/CPMP Note for guidance on plasma-derived medicinal products (CPWP/BWP/269/95, rev.3)
- ¹⁹ EMEA/CHMP Points to consider on Xenogeneic Cell Therapy Medicinal Products (CPMP/1199/02)
- ²⁰ EMEA/CHMP Note for Guidance on Use of Bovine Serum in the Manufacture of Human Biological Medicinal Product (CPMP/BWP/1793/02)
- ²¹ Eudralex Vol. 7BIm10a Table of extraneous agents to be tested for in relation to the general and species specific guidelines on production and control of mammalian veterinary vaccines
- ²² EMEA/CPMP Note for Guidance on Production and Quality Control of Animal Immunoglobulins and Immunoserum for Human use (CPMP/BWP/3354/99)
- ²³ EMEA/CHMP Note for Guidance on the quality, preclinical and clinical aspects of gene transfer medicinal products (CPMP/BWP/3088/99)
- ²⁴ Council Directive 93/42/EEC of 14 June 1993 concerning Medical Devices
- ²⁵ Council Directive 90/385/EEC of 20 June 1990 on the approximation of the laws of Member States relating to Active Implantable Medical Devices
- ²⁶ EN/ISO 10993-18:2005 Biological evaluation of medical devices- Part 18: Chemical characterization of materials
- ²⁷ EN/ISO 10993-19:2006 Biological evaluation of medical devices- Part 19: Physico-chemical, morphological and topographical characterization of materials
- ²⁸ Ph.Eur. Text 5.1.6: Alternative methods for control of microbiological quality (01/2008:50106) and General Method 2.6.27: Microbiological control of cellular products.
- ²⁹ ICH Q6B Note For Guidance on Specifications: Test Procedures and Acceptance Criteria for Biotechnological/Biological Products. (CPMP/ICH/365/96)
- ³⁰ EMEA/CHMP guideline on potency testing of cell-based immunotherapy medicinal products for the treatment of cancer (CHMP/BWP/271475/06).
- ³¹ Ph. Eur. General Text 5.2.3: Cell substrates for the production of vaccines for human use (01/2008:50203)
- ³² EMEA/CPWP Note for Guidance on Development Pharmaceuticals for Biotechnological and Biological Products (CPMP/BWP/328/99)

³³ EN/ISO 10993-1, Biological evaluation of medical devices - Part 1: Evaluation and testing

³⁴ ICH Q5E, Comparability of Biotechnological/Biological Products (CPMP/ICH/5721/03)

³⁵ ICH S6; Preclinical safety evaluation of biotechnology derived products (CPMP/ICH/302/95)

³⁶ ICH S7A, Safety pharmacology studies for human pharmaceuticals (CPMP/ICH/529/00)

³⁷ Directive 2001/20/EC of the European Parliament and Council of 4 April 2001 on the approximation of the laws, regulations and administrative provisions of the Member States relating to the implementation of good clinical practice in the conduct of clinical trials on medicinal products for human use.