Nutrition/Interventional - AUSTRIA

Competent authority

Contact Details

Contact Name 1

Bundesamt für Sicherheit im Gesundheitswesen BASG/ Federal Office for Safety in Health Care

Contact Name 2

Agentur für Gesundheit und Ernährungssicherheit AGES/ Austrian Medicines and Medical Devices Agency

Phone

0043 50 555-36111, -36820

Email Department

clinicaltrials@ages.at

Address

Traisengasse 5

ZIP/City

1200 Vienna

Country

Austria (AT)

Web address

http://www.ages.at/ages/geschaeftsfelder/medizinmarktaufsicht/

Trial Authorisation / Registration / Notification

Regulatory and ethics bodies involved in approval process

Medicines Agency Institutional Competent Authority Institutional Ethics Committee Not validated

Regulatory and ethics bodies involved in approval process for trials including patients

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Regulatory and ethics bodies involved in approval process for trials including including healthy participants

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Regulatory and ethics bodies involved in approval process for trials including vulnerable population

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Regulatory and ethics bodies involved - Additional information

Institutional competent authority is not financially involved in the approval process

CA - Registration/ notification without approval required for

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CA - Registration requirements for clinical trials

Registration recommended

Registration requirements for clinical trials including patients Registration requirements for clinical trials including healthy participants Registration requirements for clinical trials including vulnerable population Registration requirements - Additional information Registration in EudraCT is not required by law (only generally referred to soft law) Employees can be required to register clinical trials (eg Medical University of Vienna - Good Scientific Practice) **CA - Submission required to** Institutional CA Studies including patients - submission required to Studies including healthy participants - submission required to Studies including vulnerable population - submission required to Submission Format Standard application form available No Standard application form only for medicinal products Website refers to Medicinal Products only. (website in German an English available) Language of Submission Language(s) of application Official national language German Language(s) of application for trials including patients Language(s) of application for trials including healthy participants Language(s) of application for trials including vulnerable population Preferred language of application **English accepted** Case by case May vary by region Documents mandatory to be in official national language

	Documents mandatory to be in local language of study site
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	Documents mandatory to be in language of the study participant
	_
Timelines Authorisation	Time to approval CA in weeks (average)
	5
	Additional Information
	35 days by law
Safety Reporting	Sponsor must declare reportable events to
	_

Ethics committee	
Contact Details	Contact Name 1
	Institutional Ethics Committee
	Contact Name 2
	Ethics Committee Mecical University of Vienna
	Phone
	+43 1 40400 21470
	Fax
	+43 1 40400 16900
	Address
	Borschkegasse 8b/E06
	ZIP/City
	1090 Wien
	Country
	Austria (AT)
	E-Mail
	ethik-kom@meduniwien.ac.at
	Web address
	http://ethikkommission.meduniwien.ac.at/
Ethical Review - General	Submission for Ethical review mandatory for
	-
	Submission of study mandatory - Additional information
	Submission of a study to the Ethics committee is regulated on institutional level.
	Submission to CA and EC to be performed in the following order
	-
Single-Centre Studies - Ethical Review	Ethical approval (favourable opinion) to be obtained from
	Institutional EC

Ethical approval (favourable opinion) for trials including patients to be obtained from Ethical approval (favourable opinion) for trials including healthy participants to be obtained from Ethical approval (favourable opinion) for trials including vulnerable population to be obtained from Multi-Centre Studies -Ethical approval (favourable opinion) required from **Ethical Review** All local ECs of participating sites EC review not required Not validated Ethical approval in trials including patients obtained from Ethical approval in trials including healthy participants obtained from Ethical approval in trials including vulnerable population obtained from Submission of **Entitled to study submission** Application Sponsor Principal Investigator Investigator Physician Dietitian Nutritionist PhD Industry National citizen Entitled to submission of trials including patients Entitled to submission of trials including healthy participants Responsible for submission of trials including vulnerable population Prerequisites for submission / approval **Additional Information** Regional differences may exist for: Sponsor, Dietitian, Nutritionist, PhD, Industry and national citizen Standard application form available Submission Format Yes Standard application form "Forum Österreich" provides an agreement but not a law (only in German)

Language of Submission	zangaage(5) or application	
	Official national language German	
	Language(s) of application for trials including patients	
	_	
	Language(s) of application for trials including healthy participants	
	_	
	Language(s) of application for trials including vulnerable population	
	_	
	Preferred language of application	
	-	
	English accepted	
	Yes May vary by region	
	Documents mandatory to be in official national language	
	-	
	Documents mandatory to be in local language of study site	
	_	
	Documents mandatory to be in language of study participant	
	_	
Timelines Ethical Review	Time in weeks from submission to positive approval (minimum)	
	4	
	Time in weeks from submission to positive approval (maximum)	
	12	
	Time in weeks from submission to positive approval (average)	
	7	
Safety Reporting	Investigator shall report SAE to	
	Sponsor Other	
	Investigator shall report SAE in trials with patients to	
	_	
	Investigator shall report SAE in trials with healthy participants to	
	_	
	Investigator shall report SAE in trials with volunteers to	
	_	
Study specific Requirements		
Study Specific Requirements		

Sponsor	Contracts with external sponsor
	Yes
Investigator	Entitled to be principal investigator
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Language of Submission Language(s) of application

Entitled to be principal investigator for trials with patients

Physician

Entitled to be principal investigator for trials with healthy participants

Each investigator May vary by region

Entitled to be principal investigator for trials with vulnerable population

Physician

Additional Information

It is only regulated for medicinal products and medical devices who may act as principal investigator.

Study Participants - Informed Consent (IC)

Standard IC form (ICF) available

Yes

Accepted format of Informed Consent (IC) form

Written consent

Accepted format of IC form for studies including patients

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Accepted format of IC form for studies including healthy participants

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Accepted format of IC form for studies including vulnerable population

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Study Participants -Vulnerable Population

Considered as vulnerable population

Children
Elderly
Pregnant women (Pregnancy)
Unconscious Persons
Incapacitated adults
People with psychiatric disorder
People with dementia
Prisoners

Vulnerable population - Additional information

It depends on the mental status if or if not elderly are considered as vulnerable population.

it is not validated if of if not lactating women are considered as vulnerable population

Regulations concerning the inclusion or exclusion available

Yes

Regulations concerning the inclusion or exclusion

federal states monitor inclusion or exclusion of vulnerable groups.

Applicable ethical regulations

Institutional National International EU directive (2001/20/EC) Not validated

	Additional Information
	AMG - Arzneimittelgesetz apply for pharmaceuticals/drug trials
Study Participants -	Reimbursement for study participants
Compensation & Reimbursement	Optional
	Reimbursement for patients
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	Reimbursement for healthy participants
	-
	Reimbursement for vulnerable population
	-
	Compensation is limited to/provided for
	Time effort Inconvenience
	Expenses arising from study participation (e.g. Travel) Depends on study population May vary by region
	Compensation for patients is limited to/provided for
	Time effort Inconvenience Expenses arising from study participation (e.g. Travel) May vary by region
	Compensation for healthy participants is limited to/provided for
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	Compensation for vulnerable population is limited to/provided for
	Time effort Inconvenience Expenses arising from study participation (e.g. Travel) May vary by region
Funding	Trials in patients financially supported by industry
	Yes
	Trials in healthy participants financially supported by industry
	No
	Trials in vulnerable population financially supported by industry
	Yes
	Funding is an issue during the approval process
	Yes
Study Participants - Recruitment & Trial	Regulations on recruitment process exist
Outcome	No
	Recruitment process
	specific for each region
	Mandatory to inform participant of clinical trial outcome
	No

Insurance

Liability insurance or alternative arrangements for damages mandatory for

Patients/Volunteers

Obligation to contract a liability insurance for trials including patients for

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Obligation to contract a liability insurance for trials including healthy participants for

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Obligation to contract a liability insurance for trials including vulnerable population for

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Name and contact insurance companies insuring clinical research

e.g. Zürich Versicherungen

Insurance fee in € value indicated as

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Insurance fee for lowest risk research - Additional Information

€ 17,00-€ 22,00 / study Participant, depending on contract negotiations

Insurance fee in € value indicated as

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Insurance fee for highest risk research - Additional Information

€ 17,00-€ 22,00 / study Participant, depending on contract negotiations

Quality Assurance/ Quality Control (QA/QC)

Regularly performed methods

Monitoring Standard Operating Procedures (SOP) Case Report Form (CRF) Depends on study population Regional differences may exist

Regularly performed methods in trials including patients

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Regularly performed methods in trials including healthy participants

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Regularly performed methods in trials including vulnerable population

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Standards concerning quality assurance and quality control exist

Yes

Regularly performed audits

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Regularly performed audits in trials including patients

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Regularly performed audits in trials including healthy participants

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Archiving & Data Management	Regularly performed audits in trials including vulnerable population — Study documents must be kept at least (in years) —
National legislation	
General Information: Applicable Legislation & Conventions	Applied regulatory conventions Declaration of Helsinki Other ethical principles for medical research (other than Declaration of Helsinki) ICH-GCP Guidelines Other guidelines for good clinical practice (other than ICH-GCP) International regulatory requirements European regulatory requirements - European Directive 2001/20/EC, 2005/28/EC National regulatory requirements Institutional regulatory requirements
	Applied regulatory conventions in studies including patients
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Applied regulatory conventions in studies including healthy participants

Applied regulatory conventions in studies including vulnerable population

Applicable national laws

Hospital Act
Data protection Act
Genetical engineering act
Medical device act
Drug act

hospital act applies only if study is performed in the hospital setting, Genetical engineering act, Medical device act and drug act only in relevant context

Applicable national laws for patients

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Applicable national laws for healthy participants

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Applicable national laws for vulnerable population

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National regulations for volunteers exist for

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Nutrition Nutrition considered as drug

No

Yes

Tissue samples permitted - Additional information

differerences in blood and tissue samples exist

Data Protection	Yes Legal framework (on safeguarding the collection, handling, recording, keeping and/or processing of any clinical trial related data and patient files) - Additional Information standard requirements regarding data protection exists
Invasive Catheters	Invasive catheters permitted Yes Additional Information trial automatically becomes MPG §40 - Medizinproduktegesetz (medical device directive)
Definition	
Interventional Study	Pefinition in national law regulated only for drug trial in AMG (Arzneimittelgesetz (drug act) §2a, paragraph 1: "systematic investigation to: 1. demonstrate or investigate effects or side effects of drugs, 2. side effects 3. determine resorption, distribution, metabolism or excretion of drugs ethical committees extend this definition to all other investigations of this type by analogy
Nutrition Study	Definition available in national law No