

1 General information

Research question	
Rationale	
Methodology	
Project timeline and planning	

2 Requested Support

Financial Support:

Yes No (if yes, insert details below)

Personnel costs Yes No

Material costs Yes No

Analytical costs Yes No

Other costs Yes No

Other costs Yes No

Other costs Yes No

Summarized value for requested support

Support by Wörwag Pharma products:

Yes No (if yes, insert details below)

Product Name

Dosage form¹

Strength²

Required units³

Packaging Details

- Market goods (original prim. & sec. packaging)
- Comparator required specify
- Placebo required
- Blinding Required (neutral prim. & sec. packaging)
- Blinding will be organized by requestor according to GMP
- Blinding is part of requested support

Comments

3 Suggested equivalent value

What is the equivalent value suggested for the support by Wörwag Pharma?

- Final study report according to ICH-E3
- Final study report according to another format *specify*
- Regular status reports/ newsletter about trial/ study progress
-
-

¹ e.g. tablet, coated tablet, capsule, injection etc.

² Strength per unit, e.g. 300mg

³ e.g. 200 packs a 100 tablets

4 Regulatory requirements and experience of the Investigator

The clinical trial/ study corresponds to the current scientific standard in terms of study design and methodology and will be conducted in accordance to all applicable legal and regulatory principles, regulations and guidelines.

Yes No

**Applicable law & guidelines
which will be followed**

Internationally valid:

Declaration of Helsinki

ICH-GCP

GPP⁴

Regional:

EU Clinical Trials Regulation 536/2014

It is planned to publish the findings of the trial/ study in an appropriate, transparent, and timely manner.

Yes No

The investigator can present a valid GCP certificate that is not older than 3 years at trial/ study start.

Yes No Not Applicable, NIS or preclinical research

⁴ Good Pharmacoepidemiology Practice

There is recent evidence (within previous 3 years) of the investigators experience in undertaking clinical research.

Yes No

Year	Title of studies	Study category
	1.	<input type="checkbox"/> Clinical Trial <input type="checkbox"/> NIS
	2.	<input type="checkbox"/> Clinical Trial <input type="checkbox"/> NIS
	3.	<input type="checkbox"/> Clinical Trial <input type="checkbox"/> NIS
	4.	<input type="checkbox"/> Clinical Trial <input type="checkbox"/> NIS
	5.	<input type="checkbox"/> Clinical Trial <input type="checkbox"/> NIS
	6.	<input type="checkbox"/> Clinical Trial <input type="checkbox"/> NIS

The investigator can present a valid license to practice medicine.

Yes No

5 Comments

6 Data protection statement

I am interested in, and therefore give my voluntary consent, that WÖRWAG Pharma GmbH & Co.KG and its local legal entities (Flugfeld-Allee 24, 71034 Böblingen, Germany, email: info@woerwagpharma.com; Tel .: +49 (0) 7031 62 04- 0) saves my personal data (namely my name, address and information about the study for which I am requesting support), as part of processing my request, and forwards it within the company to employees of Wörwag Pharma involved in processing my request.

I am aware that I can revoke my declaration of consent (s) at any time without affecting the legality of the processing carried out based on the consent until the revocation.

Yes No

Date, full name