

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 form ¹	
Strength ²	
Required units ³	
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

- ² Strength per unit, e.g. 300mg
- ³ e.g. 200 packs a 100 tablets

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



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
	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.



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.

□ Ye	s 🔲 No	
Year	Title of studies	Study category
	1.	☐ Clinical Trial ☐ NIS
	2.	Clinical Trial NIS
	3.	Clinical Trial NIS
	4.	Clinical Trial NIS
	5.	Clinical Trial NIS
	6.	☐ Clinical Trial ☐ 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