

Course Specifications

Valid as from the academic year 2024-2025

Pharmaceuticals in the Environment (J000529)

Course size Credits 6.0		(nominal values; actual values may depend on programme) Study time 150 h				
Course offerings and	teaching methods in academic					
A (semester 1)	English	Gent	p	oractical		
			i	ndependent wor	k	
			g	roup work		
			e	excursion		
			S	eminar		
			l	ecture		
Lecturers in academi	c year 2024-2025					
Hertleer, Carla			FW02	staff membe	er	
Wynendaele, Evelien FWO2			FW02	lecturer-in-charge		
De Spiegeleer, I	Bart		FW02	co-lecturer		
Offered in the following programmes in 2024-2025				crdts	of	
International Master of Science in Sustainable Drug Discovery				6		
Teaching languages						
English						
Keywords						
•	narmaceuticals; eco-toxicity; risk a	ssessment				
Position of the cours						
Different concer pharmaceutical are provided wi of pharmaceutic out by the stude animal and plar	tudents are introduced into the en ots and principles of environmenta s and related health products are th extensive knowledge about env cals. Theoretical principles are illus ents as well as practical laboratory at eco-toxicity of medicines, pharm tical waste treatment will be turn	al sustainability related elucidated. Moreover, s vironmental risk assess strated by case studies y sessions and visits: co naceutical environmen	d to students sment (ERA) s worked- oncepts of			
Contents						
	hazards and risks, efficacy and sa iversity, one health, cradle-to-crac		ıd eco-			

pollution/contaminants, green chemistry, compartments, systems thinking
2) Environmental consequences of pharmaceuticals: levels (order of magnitude), different medicine classes, different elements of a medicine (active pharmaceutical ingredient, excipients, packaging)

3) Eco-toxicity tests: experimental design, experiments with *C. elegans* (icw prof. Braeckman), plants (icw prof. Werbrouck) and lab-analyses, dose-response/effect analysis

4) Exposure analysis: approaches, uncertainties

5) Environmental Risk Assessment (ERA) of medicines: models, EU-guidelines (human, veterinary)

6) Mitigation/reduction, incl. waste treatments (incl. plant visit) Benign-by-Design (BbD) in drug discovery

Initial competences

0.0h 0.0h 0.0h 0.0h 0.0h

offering A Basic knowledge of fundamental physical, chemical and biological principles and processes.

Final competences

- 1 Understand semantics
- 2 Explain systems thinking and the place of pharmaceuticals in the earth's ecosystem
- 3 Perform a conceptual ERA analysis of a pharmaceutical
- 4 Discuss balance of efficacy, safety, quality versus ecotoxicity of medicines
- 5 Have insights in the used techniques and methods in the ERA of pharmaceuticals
- 6 Understand the applied modelling & probabilistic approaches
- 7 Discuss different approaches to reduce the environmental impact of pharmaceuticals
- 8 Demonstrate the ability to understand, summarize and evaluate relevant scientific literature
- 9 Communicate scientific rationales and critical reflections leading to concise conclusions

Conditions for credit contract

Access to this course unit via a credit contract is determined after successful competences assessment

Conditions for exam contract

This course unit cannot be taken via an exam contract

Teaching methods

Group work, Seminar, Excursion, Lecture, Practical, Independent work

Extra information on the teaching methods

Eight (8) lectures of each 1.25 hours will be given. Four (4) coached seminars of 1.25 hours each, plus two (2) guided self-study sessions (each 1.25 hours) for the oral presentation. 1 study visit (1 day) on the waste treatment, mitigations and adaptations of a pharmaceutical company. Eighteen (18) practical sessions (each 1.25 hours) are given, divided in different topics (incl. eco-toxicity, pharmaceutical analyses and planttoxicity/bioremediation). Three (3) additional coached seminar sessions (each 1.25 hours) are organised with exercises about read-across systems.

Study material

Type: Slides

Name: Slides Indicative price: Free or paid by faculty Optional: no Available on Ufora : Yes

References

Is available in course material.

Course content-related study coaching

Guidance of students in developing a presentation on an ERA of a pharmaceutical.

Assessment moments

end-of-term and continuous assessment

Examination methods in case of periodic assessment during the first examination period

Oral assessment, Written assessment

Examination methods in case of periodic assessment during the second examination period

Oral assessment, Written assessment

Examination methods in case of permanent assessment

Participation, Peer and/or self assessment, Assignment

Possibilities of retake in case of permanent assessment

examination during the second examination period is possible in modified form

Extra information on the examination methods

Periodic evaluation: oral exam starts after a written preparation.

Calculation of the examination mark

The seminar activities, leading to an oral presentation (in the last coached seminar for oral presentation), and the practical sessions, leading to different reports, are responsible for 50% of the course score. The examination at the end of the semester is responsible for the other 50% of the points. If the student obtains a score below 9/20 for one of the separate parts of the evaluation (seminar activities, practical sessions or examination), then the final mark is automatically reduced to the lowest score obtained for one of the separate parts. *Students who eschew period aligned and/or non-period aligned activities or evaluations in this course may be failed by the examiner.*