

Subject card

Subject name and code	Aspects of implementation and certification of medical products, PG_00053349								
Field of study	Biomedical Engineering, Biomedical Engineering, Biomedical Engineering								
Date of commencement of studies	February 2025		Academic year of realisation of subject		2025/2026				
Education level	second-cycle studies		Subject group		Optional subject group Specialty subject group Subject group related to scientific research in the field of study				
Mode of study	Full-time studies		Mode of delivery			at the university			
Year of study	2		Language of instruction		Polish polski				
Semester of study	3		ECTS credits		2.0				
Learning profile	general academic profile		Assessment form		assessment				
Conducting unit	Department of Biomedical Engineering -> Faculty of Electronics, Telecommunications and Informatics								
Name and surname	Subject supervisor		prof. dr hab. inż. Jerzy Wtorek						
of lecturer (lecturers)	Teachers		prof. dr hab. inż. Jerzy Wtorek						
Lesson types and methods of instruction	Lesson type	Lecture	Tutorial	Laboratory	Projec	roject Seminar		SUM	
	Number of study hours	15.0	15.0	0.0	0.0		0.0	30	
	E-learning hours included: 0.0								
Learning activity and number of study hours					Self-study S		SUM		
	Number of study hours	30		2.0		18.0		50	
Subject objectives	The subject introduce regulations regarding							g from the	

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Learning outcomes Course outcome		Subject outcome	Method of verification				
	[K7_U71] is able to apply knowledge from humanistic, social, economic or legal sciences in order to solve problems	The student is able to design clinical trials taking into account the aspect of their credibility, taking into account the actual conditions occurring during these trials. The student is able to select legal requirements relating to the certification process.	[SU4] Assessment of ability to use methods and tools				
	[K7_W10] knows and understands, to an increased extent, the basic processes occurring in the life cycle of equipment, objects and technical systems, as well as methods of supporting processes and functions, specific to the field of study	Based on the knowledge obtained, the student knows and understands the principles and requirements for the correct process of designing, implementing, putting into service and monitoring a medical device.	[SW3] Assessment of knowledge contained in written work and projects [SW1] Assessment of factual knowledge				
	[K7_W11] knows and understands, to an increased extent, the general principles of creation and development of forms of individual entrepreneurship and the economic, legal and other conditions of various types of activities related to the awarded qualification, including the principles of protection of industrial property and copyright law	The student knows the regulations governing the admission of medical products to the market and the related requirements for the design, production and quality assurance management system.	[SW1] Assessment of factual knowledge				
	[K7_K71] is able to explain the need to apply knowledge from humanistic, social, economic or legal sciences in order to function in a social environment	Based on the acquired knowledge, the student understands the need to conduct properly designed clinical and market research and the need to use management systems that take into account appropriate legal regulations.	[SK5] Assessment of ability to solve problems that arise in practice				
Subject contents	Discussion of the design, implementation and certification cycle of a medical product.2. Importance and basic requirements for the prototype, MVP and final product.3. Certification process.4. Clinical trials at various stages of product development.5. Designing clinical trials.6. Market research and obtaining funds for various stages of medical product development.						
Prerequisites and co-requisites	Basics of electronics, physics and st	atistical methods in medical applicati	ions				
Assessment methods	Subject passing criteria	Passing threshold	Percentage of the final grade				
and criteria		55.0%	40.0%				
	umiejętności	55.0%	60.0%				
Recommended reading	Basic literature Medical Devices Regulation (MDR) ISO 14155 Clinical Investigation of Medical Device Methodology for the clinical development of medical devices, French National Authority for Health						
	Supplementary literature	https://single-market-economy.ec.europa.eu/sectors/electrical-and-electronic-engineering-industries-eei/radio-equipment-directive-red_en https://eur-lex.europa.eu/legal-content/PL/TXT/? uri=celex%3A32014L0035					
	eResources addresses	Adragy na platformia oNaugrapio:					
	ersesources addresses	Adresy na platformie eNauczanie:					

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Example issues/ example questions/ tasks being completed	Discuss the concept of MVP2. Based on the design documentation, carry out a simulated device certification procedure.3. Principles and requirements for designing clinical trials.
Work placement	Not applicable

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