		築品・医療機器の開発計画、薬事と審査 lopment strategy, plan, and regulatory affairs of drugs and medical devices				de	filiated partment b title,Na	י ה		duate School of Medicine fessor,KAWAKAMI KOJI	
Grade allot	əd	Professional degree	students	Number	of cred	its	2		se offered period	2016/Second semester	
Day/period	1	Wed.3,4	Cla	ss style	Lecture	e			Language	Japanese and English	
[Outline and Purpose of the Course] Director and Instructors:											
 Cardiovascular Center), Hiromu Kutsumi (Shiga Univ. Of Medical Science), Ikuo Horii (Cambridge Univ.), Yoko Uryuhara (Doshisya Univ.), Shigeyuki Wakitani (Mukogawa Women's Univ.), Hisashi Urushihara (Keio Univ.), Shinya Kimura (Japan Medical Data Center), Yoshie Onishi (CreativCeutical) The development strategy, protocol design, project management, safety/efficacy assessment, and economic evaluation of drugs and medical devices will be lectured. The development of drug and medical device and the regulatory review in terms of manufacturing and control, nonclinical studies, clinical protocol, the new drug applications, and post-marketing will be discussed. [Course Goals] The idea of drug/medical device development and regulatory review will be understood. To understand the strategy, protocol development, and project management of drug development and clinical trials. 											
[Course Schedule and Contents] Lectures and discussions											
 Lecture sch October 5 October 12 October 19 October 26 November 2 November 9 November 16 November 30 December 7 December 14 December 21 January 4 January 11 January 18 	ied Dru nd nd nd Pr Gl Ii Ii Da I I Ir ndu Apj Hea	ule 1g Developme ustry R&D an ustry R&D an	d regu d regu nent o nent S and re and re perso agement ind regu l regu edical and ou	ulatory revi ulatory revi ulatory revi of clinical the strategy of 2 gulatory re- onalized me ent of drug gulatory revi latory revie database in utcomes res	ewer tra ewer tra ewer tra trial (Niin New Dru viewer tr dicine in (Uryuha viewer train wer train evaluat earch: P	inin inin inin mi) ugs rain rain rain rain rain rain rain rain	g: nonci g: CMC g: proto (Elze) ing: me ing: me ug deve ing: regu g: postm of healt ice (On	linical C (Kaw col re dical c dical c lopme enerat arket h care	studies (Kor vakami) view of clinic levices (1) (V levices (2) (K nt (Horii) ive medicine (Urushihara)	cal trial (Fujiwara) Vamamoto) Kutsumi)	
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医薬品・医療機器の開発計画、薬事と審査(2)

[Class requirement]

You must take this course along with the course H074 "Drug policy and regulations" and H079 "Drug development, evaluation and regulatory sciences" of Wednesday 2nd of the second semester for you to acquire fundamental understanding of the field.

[Method, Point of view, and Attainment levels of Evaluation]

Participation (50%) and report (50%)

[Textbook]

Suggested reading:

Saeko Yasuo et al. Invitation to new drug development. Kyoritsu Press, 2006 (Japanese). Koji Kawakami (ed.) The Beginner's guidance to the clinical research and development. Medical Do, 2010 (Japanese).

[Reference books, etc.]

(Reference books) Introduced during class

[Regarding studies out of class (preparation and review)]

Preparation in advance and review after lecture

(Others (office hour, etc.))

Koji Kawakami (3F, Bldg G) appointment required by email. e-mail: kawakami.koji.4e@kyoto-u.ac.jp

*Please visit KULASIS to find out about office hours.