

MODULE DESCRIPTOR

Module Title

Drug Dosage Form and Design

Reference ASM041 Version 4 Created October 2022 SCQF Level SCQF 11 **SCQF** Points Approved May 2019 30 Amended October 2022 **ECTS Points** 15

Aims of Module

To enable the student to develop a detailed understanding of the design of dosage forms, taking into account the preclinical and clinical aspects of medicine development, within a quality framework and in the context of a variety of clinical indications and patient groups.

Learning Outcomes for Module

On completion of this module, students are expected to be able to:

- 1 Demonstrate a critical understanding of the application of pharmaceutics in the design of dosage forms.
- Have a detailed understanding of the pre-clinical and clinical development process required and the underpinning quality framework which supports such development.
- 3 Demonstrate a critical understanding of the biopharmaceutical concepts surrounding the development of medicines.

Indicative Module Content

Topics include the processes underpinning the development of dosage forms such as liquids, suspensions, emulsions, topicals, parenterals, suppositories, tablets and capsules. In addition the pre-clinical and clinical considerations of medicine development will be covered including preclinical testing (an overview of ADME and DMPK studies) and the stages of clinical trials, together with aspects of quality systems which apply to dosage form development, including cGMP, GLP and GCP. The biopharmaceutical basis for the development of medicines including the Noyes-Whitney equation and how physiology affects drug absorption, physicochemical and dosage form factors which affect bioavailability and dosage regimes for a range of patients at all stages of life from paediatrics to geriatrics.

Module Delivery

The module will be delivered using lectures and coursework supported with tutorials.

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Indicative Student Workload	Full Time	Part Time
Contact Hours	76	N/A
Non-Contact Hours	224	N/A
Placement/Work-Based Learning Experience [Notional] Hours	N/A	N/A
TOTAL	300	N/A
Actual Placement hours for professional, statutory or regulatory body		

ASSESSMENT PLAN

If a major/minor model is used and box is ticked, % weightings below are indicative only.

Component 1

Type: Practical Exam Weighting: 50% Outcomes Assessed: 1

Description: A presentation of the development of medicines laboratory work.

Component 2

Type: Coursework Weighting: 50% Outcomes Assessed: 2, 3

Description: Series of essays throughout the semester. The grade awarded is an average of all essays

undertaken.

MODULE PERFORMANCE DESCRIPTOR

Explanatory Text

The first grade represents Component 1 (CW1) and the second Component 2 (CW2) both of which are equally weighted. A minimum module grade of D is required for a pass, with compensation of grade E in either CW1 or CW2 permitted. Non-submission of either component will result in an NS grade.

The pointing of the control of the component will recall in all the grade.		
Module Grade	Minimum Requirements to achieve Module Grade:	
Α	AA,AB,BA	
В	AC,AD,BB,BC,CA,CB,DA	
С	AE,BD,BE,CC,CD,DB,DC,EA,EB	
D	CD,DD,DE,EC,ED	
E	AF,BF,CF,DF,EF,FA,FB,FC,FD,FE	
F	FF	
NS	Non-submission of work by published deadline or non-attendance for examination	

Module Requirements

Prerequisites for Module None, in addition to course entry requirements.

Corequisites for module None.

Precluded Modules None.

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INDICATIVE BIBLIOGRAPHY

Aulton, M.E.(ed.), Taylor, K. (2017)'Aulton's pharmaceutics - the design and manufacture of medicines', 5th edition, Churchill Livingstone, Elsevier.

- 2 Jones, D. (2016). 'Pharmaceutics dosage form and design'. Fasttrack, 2nd edition, Pharmaceutical Press.
- 3 Denton, P., Rostron, C. (2013). 'Pharmaceutics the science of medicine design', Oxford University Press.