8 November 2021, Strasbourg, France

PDG signs-off on milestone harmonised general chapter on chromatography

The harmonised general chapter *Chromatography* was signed-off by the Pharmacopoeial Discussion Group (PDG), which brings together the European Pharmacopoeia (Ph. Eur.), Japanese Pharmacopoeia (JP) and the United States Pharmacopeia (USP), on 28 September 2021. The coordinating pharmacopoeia for this text was the Ph. Eur.

During a joint PDG–industry meeting in 2009, the PDG was encouraged to add harmonisation of the three regional chapters on chromatography to the PDG work programme. Although the chapters in question differed in content and format, it was considered feasible to develop a chapter describing *core requirements* applicable for TLC, HPLC and GC.

After discussion, it was agreed not to include more general (textbook type) descriptions of individual techniques as each of the PDG pharmacopoeias has its own approach, decided at regional level.

These harmonised requirements promote the development of individual monographs with a consistent approach and enhance understanding of basic requirements by users in all three regions.

Owing to the complexity of the topic, the differences in the individual approaches mentioned above and the large impact on existing monographs or general chapters, considerable time and effort were invested in finding the best possible compromise.

During harmonisation of the text, particular attention was paid to:

1. Terminology, definitions and interpretation of chromatograms;
2. System suitability: this section provides requirements intended to guarantee that the performance of the chromatographic system is appropriate. They apply to multiple monographs and are to be read in conjunction with the requirements described therein;
3. Adjustment of chromatographic conditions;
4. Quantitation procedures.

The corresponding regional texts are scheduled for publication in July 2022 (Ph. Eur.), December 2022 (JP) and December 2022 (USP).

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**Note for the Editor:** Further information is available on the internet site https://www.edqm.eu/.

The EDQM is a leading organisation that protects public health by enabling the development, supporting the implementation and monitoring the application of quality standards for safe medicines and their safe use. Its standards are recognised as a scientific benchmark worldwide. The European Pharmacopoeia is legally binding in member states. The EDQM also develops guidance and standards in the areas of blood transfusion, organ transplantation and consumer health issues.
1. There are 40 members of the European Pharmacopoeia Commission: Austria, Albania, Belgium, Bosnia and Herzegovina, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Republic of Moldova, Montenegro, Netherlands, North Macedonia, Norway, Poland, Portugal, Romania, Serbia, Slovak Republic, Slovenia, Spain, Sweden, Switzerland, Turkey, Ukraine, United Kingdom and the European Union.

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