Regulatory Science and Innovation Programme for Europe (ReSciPE): a proposed model

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Abstract

Regulatory science underpins the objective evaluation of medicinal products. It is therefore imperative that regulatory science and expertise remain at the cutting-edge so that innovations of ever-increasing complexity are translated safely and swiftly into effective, high-quality therapies. We undertook a comprehensive examination of the evolution of science and technology impacting on medicinal product evaluation over the next 5-10 years and this horizon-scanning activity was complemented by extensive stakeholder interviews, resulting in a number of significant recommendations. Highlighted in particular was the need for expertise and regulatory science research to fill knowledge gaps in both more fundamental, longer-term research, and with respect to technological and product-specific challenges. A model is proposed to realise these objectives in Europe, comprising a synergistic relationship between the European Medicines Agency (EMA), the European Medicines Regulatory Network and academic research centres to establish a novel regulatory science and innovation platform.

What is already known about this subject?

The EMA and medicinal product regulators around the world, are confronted continuously with advances in science and technology. However, the complexity of innovation is increasing rapidly, requiring regulatory science to evolve in tandem and to develop an effective mechanism to do so in a timely manner.

What this study adds?

This study explores regulatory science needs over the next 5-10 years and proposes a mechanism to enable regulatory science to keep pace with innovation.
Introduction

Translating fundamental science into patient-accessible therapies requires application of diverse scientific disciplines. Regulatory science underpins the objective evaluation of the safety, efficacy and quality of medicinal products and crucially informs the regulatory decision-making process.

Specifically, therefore, regulatory science must provide medicines’ regulators with the knowledge to apply innovative research and novel methodological tools to the objective determination of the benefits and risks associated with the use of a new medicinal product\(^1\). It is fair to say, however, that rapid progress in the biomedical and related sciences – for example, in areas such as cell-based therapies, drug-device combinations, predictive toxicology and artificial intelligence – mean that the most challenging regulatory questions\(^2 - 4\) are originating from the fastest moving and most competitive scientific disciplines\(^5\). As a result, it is absolutely imperative that regulatory science remains at the cutting edge so that innovations of ever-increasing complexity are translated safely into efficacious and affordable therapies in a timely fashion, promoting public health.

The European Medicines Agency (EMA) engages continuously with advances in regulatory science and, in 2017, undertook a comprehensive baseline review examining the evolution of science and technology that will impact its core business of medicinal product evaluation over the next 5-10 years. This horizon-scanning activity was complemented by an extensive stakeholder outreach exercise across individuals and organisations involved in the entire medicine development lifecycle (and included, *inter alia*, the pharmaceutical industry, health technology assessors and payers, regulatory science experts, academia, scientific organisations and societies, European Union research infrastructure networks, healthcare professionals and patient representative groups). The cumulative result of this concerted effort was a document\(^6\), “EMA Regulatory Science to 2025 – Strategic Reflection”, currently released for public consultation at the end of 2018 and recently summarised in the literature\(^7\). A key component of this reflection is a proposed model to strengthen regulatory science and innovation in Europe, the elaboration of which is now described\(^6\).
**Methods**

1. **Horizon scan (baseline review)**

   The initial (>60) areas of review (see Supplementary Information, Table S1) across health, science, technology and regulatory science were selected by the EMA’s internal scientific leadership, the Scientific Coordination Group (SCG). Subsequently, a multidisciplinary research group conducted an initial horizon scanning exercise. This included mining, *inter alia*, internal databases and the relevant scientific literature. In each area reviewed, the state-of-play and the projected opportunities and challenges over the coming 5-10 years were identified. These results were authenticated within the research group, and then peer-reviewed by in-house experts and the SCG.

2. **Stakeholder interviews**

   Interviews were then carried out with external experts and key opinion leaders from the EMA’s principal stakeholder groups to validate the internal conclusions. Interviewees were nominated by the European Medicines Regulatory Network (EMRN) and drawn from the Agency’s expert database; non-response error was mitigated through follow-up reminders. The interviews (n = 70) were either semi-structured (55) or open (15). The stakeholders were provided with a series of key questions (developed by the research group) and an introduction to the baseline review prior to the interviews. The questions were aligned with the aims of the regulatory science reflection and were trialled with colleagues, and re-ordered and optimised in terms of timing. The resultant draft script was then tested on an initial panel of interviewees for feedback. This feedback was incorporated into a final master script targeted towards semi-structured interviews with each stakeholder group. For the open interviews, the script was used after the interviewees had provided their unprompted, initial topics for discussion.

3. **Data acquisition and analysis**

   The semi-structured interviews lasted around 1 hour, the open interviews up to 2 hours. A written record of the interviews was made by two or more of the research team and then cross-checked for accuracy and consistency. Analysis of the information obtained involved open and axial coding where the research team attributed codes to meaningful sections of text (words, statements and sentences). These codes were compared and a subset agreed before undertaking additional rounds of axial coding. The findings were eventually reported using Consolidated Criteria for Reporting Qualitative Research (COREQ). Finally, the codes were grouped into themes, which were compared to and merged with the results of the horizon scan and baseline review. From this exercise, a set of overarching strategic goals for regulatory science emerged along with a number of core recommendations and associated underlying actions necessary to achieve these aims.
Results

The baseline review, horizon scan and stakeholder outreach resulted in over 600 comments and recommendations. Many of these identified the need for expertise and regulatory science research to fill knowledge gaps in two broad areas as discussed in detail in the published EMA document, "EMA Regulatory Science to 2025 – Strategic reflection" and summarised elsewhere: (i) those requiring more fundamental, longer-term research, and (ii) where technology or product-specific challenges were evident. Relatedly, the limited funds available for regulatory science research, and the clear need for more resource in this area, represented very strong signals.

Regarding expertise, a deficit in the area of regulatory science know-how was identified, particularly in rapidly evolving domains of research and innovation such as drug-device combinations, predictive toxicology and artificial intelligence. A more proportionate approach to access international expertise was a recurring suggestion in this regard. Enhanced training in the relevant science for stakeholders and regulators alike was also highlighted.
Discussion

The primary role of medicines regulatory agencies may be summarised as one of protecting and promoting public health and, increasingly, by catalysing and enabling science to be translated into patient-centred healthcare\(^1\). To meet these objectives, the regulatory agency must understand the fundamentals of the relevant science, and their application in the medicinal product review and approval process, and be critically informed of key areas of scientific innovation that have the potential to impact on its core business\(^5,6\).

A model to underpin regulatory science and innovation in Europe

A mechanism with which these goals can be achieved in Europe is a synergistic relationship between the EMA, the EMRN and distributed academic research centres to establish a novel science and innovation platform – provisionally termed the Regulatory Science and Innovation Programme for Europe (ReScIPE) – that undertakes both long-term, fundamental research in strategic areas of regulatory science (Figure 1, upper panel), and shorter-term investigations to address emerging regulatory science questions (Figure 1, lower panel).

**Figure 1**: Upper panel - An iterative partnership between regulators, European public funding agencies and academic scientists to strategically focus basic research in regulatory science. The potential funding agencies include those at the European level, such as DG RTD and IMI, and national funders.

Lower panel - Research collaboration between network scientists and academia to tackle rapidly-evolving regulatory science questions and to translate innovation efficiently into regulatory tools and processes.

**ReScIPE: goals and deliverables**

It is anticipated that ReScIPE will identify research priorities that promote the field of regulatory science - including innovative research, development of regulatory tools, education, and scientific
exchange - together with not-for-profit and commercial entities striving to produce safe, effective, affordable and high-quality medical products. Self-evidently, collaboration involving ReSciPE and the European pharmaceutical, biotechnology, and high-tech industries is particularly important to the long-term aims articulated above. With the governance of these collaborations being carefully decided by funders at the call stage. It is also envisaged that partnerships between EMA, the EMRN and academia will also develop regulatory training modules and undertake horizon scanning in emerging areas of innovation, and that ReSciPE will drive a data-sharing culture to foster open science that is mutually beneficial for all stakeholders.

**Precedence for success in Europe**

Given the strength of the pharmaceutical and biotechnology industries in Europe, the established importance of leading scientific professional societies (such as EUFEPS, the European Federation for Pharmaceutical Scientists), the considerable regulatory expertise at EMA and across the EMRN, and the world-leading quality of biomedical research related to medical product innovation and development in European universities and research centres, the present situation also affords a real opportunity to accomplish a paradigm-shift in regulatory science and innovation through the establishment of ReSciPE. This concept must build upon precedents at the national level, including the Dutch Medicines Evaluation Board (MEB) Regulatory Science Program, which has led to the creation of a broad network of partnerships between academic and other external parties. In this way, MEB has committed a budget to catalyse and facilitate both short-term projects and longer-term PhD theses to enhance its ability to deliver high quality benefit/risk assessment. Three specific areas of the medicinal product lifecycle have been targeted: development and innovation, regulation and decision-making, and consumer use and safety. At the same time, MEB is actively participating in regulatory education and learning, for example, via internships to bachelor- and masters-level students. Other similar research models include Germany’s Federal Institute for Drugs and Medical Devices (BfArM), which conducts research in collaboration with national, EU and international research centres and academia, and the Paul-Ehrlich-Institut, (PEI), which interacts with leading research institutes, academia and international organisations to set new standards in the field of vaccines/biomedicines. Another example is the European Center of Pharmaceutical Medicine (ECPM), based at the University of Basel, that provides training which covers the entire medicinal product development process from molecule identification to commercialisation, including an understanding of essential aspects of regulatory science.

Most recently, a new EU-funded project entitled “Strengthening Training of Academia in Regulatory Science” (STARS), was initiated. The consortium involved includes the EMA and 20 regulatory bodies. The three-year project aims to analyse and improve the training of academia in regulatory science and to enhance regulatory protocol assistance in academic-driven health research. These measures are designed to facilitate translational clinical research in academia, and to accelerate the availability of innovative, cutting-edge therapies to patients across Europe.

**CERSIs: an American model**

Furthermore, evidence from the US, in particular, suggests that this model of synergistic partnership between a regulatory agency, academic researchers and key stakeholders, such as established pharmaceutical companies and small and/or medium-sized enterprises, is a fruitful approach to ensure that research ideas are effectively translated into new and effective medical products and that
technological advances resulting in novel tools are applied to catalysing and facilitating the regulatory
review and approval process, thereby accelerating patient access to innovative therapies\textsuperscript{16}. The US
Food & Drug Administration (FDA) currently funds five Centres of Excellence in Regulatory Science &
Innovation, each with a particular focus associated with the Agency’s priority areas\textsuperscript{17}. The UCSF-
Stanford Centre, for example, is addressing the over-arching strategic aim to develop new models and
methods for moving drugs and other medical products, such as devices and cell-based therapies, from
the laboratory to clinical trials\textsuperscript{18}. In parallel, the Centre provides training and educational programs
(including internships and laboratory rotations) for PhD students, postdoctoral fellows, faculty and
scientists in the industry and at the FDA.

\textbf{Conclusions}

Scientific challenges in regulatory science and innovation span the entire spectrum of the medicinal
product lifecycle -- for both human and veterinary drug product development\textsuperscript{19} -- from, for example,
the conception and development of new cell-based treatments, through new thinking in predictive
toxicology, and the rapidly increasing variety of imaginative drug-device combination products, to new
ideas concerning the personalisation and precision of medical therapy (including the manufacturing
challenges)\textsuperscript{5,6}. As such, there is a strong rationale for ReSciPE to use a distributed model, and to
benefit from the collaboration of expertise across different academic centres that each concentrate
on specific target areas of investigation.

The scale of investment required is logically a function of the number and complexity of the
transformational research questions to be addressed, the requirements for associated infrastructure,
and the perspective taken on the specific role of ReSciPE in training early-career scientists in this
important field. In developing existing interactions between the EMA, the EMRN and academia (as
well as integrating with ongoing key European activities as mentioned above) to ensure that
regulatory science keeps up-to-date, these resources must also be proportional to the public health
aim of ensuring that medicines’ regulation not only guarantees safe and effective therapies that meet
the highest standards of quality, but that it also facilitates patient access to these innovative and
important medicines\textsuperscript{5}. While this latter challenge is one with which regulators are wrestling to an
ever-increasing extent, further discussion of how to achieve better and more uniform access to novel
(and almost always expensive) therapies, and to a high standard of healthcare in general, is beyond
the scope of this article.

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