What next for US clinical trials?

Part 5 of MESM’s series on proposed federal research regulation amendments and the impacts on commercial sponsors and research organisations

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Introduction

In part one of this series, we introduced a broad overview of the proposals and proposed timelines for changes to the Common Rule of HHS Regulation (45 CFR 46, Subpart A) and the Notice of Proposed Rule Making (NPRM).

In part two, we discussed the reclassification of biospecimens as human subjects under the NPRM and the potential impacts of these proposed changes to clinical researchers.

In part three, we reviewed the proposed changes to informed consent and the potential impacts for life science companies and academic research institutions.

What happened during the NPRM consultation process?

A consultation period inviting comments from industry and other stakeholders was extended to the 6th January 2016 following the publication of the proposed NPRM in September 2015.

During that period the OHRP received over 2,100 comments, of which a majority raised concerns over the proposed provisions for the treatment of biospecimens and informed consent. Many comments also offered potential solutions to foreseen problems.

A consultation period gave industry stakeholders the chance to comment on the proposed changes.

Whilst scientists were generally supportive of the need for an update to the Common Rule to take account of recent advances in science and research, there was broad concern that some of the changes would inhibit or slow research considerably, and that the proposed changes were unduly complicated and/or onerous.
A sample of concerns raised during commenting

The Infectious Diseases Society of America (ISDA) raised concerns that requiring informed consent for all human specimens, including those that had been de-identified “would have a chilling effect on many types of research”; and argued that the logistics of implementation were too daunting and unrealistic for most research organisations. Instead they suggested there should be strong penalties against re-identification of biospecimens.1

Similar concerns were also expressed in comments from Stanford University. In addition, a particular concern raised was that health care facilities that serve minorities and economically disadvantaged populations would not have the financial resources available to obtain and track consent, with a consequence that research would be increasingly performed with biospecimens from a skewed population.2 As such, research findings may no longer be applicable to all population groups.

The Advanced Medical Technology Association (AdvaMed) expressed concerns that costs of research would be raised 10 to 50 fold, and that studies that previously took weeks to conduct would require months or years. AdvaMed stated that this “would be starkly inconsistent with FDA's statutory and regulatory mission of promoting public health and ensuring that medical devices provide a reasonable assurance of safety and effectiveness.”3

The Pharmaceutical Research and Manufacturers of America (PhRMA) commented that researchers and hospital administrators overseeing biorepositories would need to develop tracking systems to match stored biospecimens along with their consent status. This would represent a considerable administrative and financial burden for these organisations.4

The American Medical Association (AMA) asked who would be responsible for gaining informed consent from patients whose specimens were not originally donated for research. I.e. whether it should be the biorepository or the requesting researcher. They expressed the concern that the administrative burden of gaining consent from such patients may lead to hospitals ceasing availability of such samples to researchers. Particular concerns were also raised by AvaMed and PhRMA in relation to genetic research to identify disease biomarkers. Such research requires access to vast stores of samples many of which may belong to patients who are terminally ill. Contacting such patients for consent could be considered to be unethical.2,4
What next for the NPRM?

The comments and suggestions received during the consultation process are now under consideration by the regulating authorities, and it is by no means clear what wording the Final Rule will contain nor how close the Final Rule will be to the original wording of the NPRM proposals. Regulators are under no obligation to change the legal wording of the Final Rule to address all comments and criticisms raised.

It is also impossible at this stage to determine whether the changes currently proposed will actually inhibit or slow down research overall if they are introduced as proposed. Some of the proposed changes will reduce research burdens whilst others will potentially create added burdens and costs.

However, it is to be expected that some form of the NPRM proposed changes encompassing more stringent guidelines for data security, informed consent and increased protection for biospecimens will go through.

If the progression to Final Rule goes ahead according to schedule, it will be published in September 2016 in the final days of President Barack Obama’s administration. However, if publication is delayed it will pass to a new administration to sign and finalise and much will depend on the priority given to this legislation under the new administration.

For now, researchers must watch, wait and see what happens next.

What next for life sciences companies?

Given the increased importance of biospecimen research to life science companies, if the NPRM is finalised in its current form it could force meaningful changes to the external and internal research practices of companies carrying out or sponsoring research during the development of new drugs, devices and other biological products.

There are three main reasons why life science companies could be impacted by the NPRM or ‘Common Rule.’

1. Whilst the NPRM will apply only to federally funded research and to ‘certain clinical trials’, life science companies frequently fund research at institutions and medical centres that also carry out federally funded research. If the NPRM changes require modifications to existing research protocols at those research institutions, then those alterations will apply to all research carried out at those centres in the future, whether they are funded by the state or industry.

2. If the NPRM proposed changes for a streamlined IRB process are introduced, life science companies will need to modify their own internal review processes in order to comply with external requirements and avoid potential delays in gaining research approvals.
3. Most commercial life science companies develop products whose developmental research falls under the FDA, whether conducted internally or by third party research institutions. The FDA have indicated that changes to the Common Rule will result in parallel changes to its own research regulations. This would necessitate substantial changes to operational systems and research protocols of all life science companies. As a result, the amendments to the Common Rule could have a greater impact on life science companies than originally anticipated.

Summary

In preparation for the proposed changes, life science companies should carefully track the progress of the NPRM and closely examine their internal organisational processes to determine whether they are robust enough to support compliance, as well as consider the potential longer term ramifications for their organisation.

Each company needs to examine these questions:

Based on current proposals, how will the NPRM affect our individual company’s research efforts if the NPRM proposals progress to Final Rule in their current form?

What will we need to change in terms of how we conduct research both internally and externally, and how will we do this?

The answers to these questions can then be reassessed on publication of the Final Rule.

References

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