Redefining Risk Stratification and Endpoints for Clinical Trials in Kidney Transplantation: Rationale and Methodology of Proposals Submitted to the European Medicines Agency by the European Society for Organ Transplantation

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The European Society for Organ Transplantation (ESOT) submitted a Broad Scientific Advice request to the European Medicines Agency (EMA) in 2018, to explore whether updating guidelines on clinical trial endpoints would encourage innovations in kidney transplantation research, thereby improving long-term outcomes for allograft recipients. The request was refined collaboratively by the EMA and ESOT, with the EMA issuing a final response in December 2020. This Transplant International special issue explores the topics that were the focus of these interactions between the EMA and ESOT. Articles explore the current issues and dilemmas in kidney transplantation, primarily relating to unclear or outdated risk stratification and markers of transplantation success, although several potential improvements for outcomes assessment are also suggested. Discussions between the EMA and ESOT and recommendations are summarized, in the hope that this project will generate further discussion eventually generating a consensus on clinical trial endpoints and risk stratification, increase the quality of research in transplantation medicine, and improve long-term outcomes for kidney transplant recipients.

Keywords: kidney transplantation outcome, EMA guideline, efficacy endpoint, long-term outcome, improvement, European Society for Organ Transplantation, risk stratification

INTRODUCTION

Over many decades, progress in the treatment of acute rejection markedly improved the short-term success of kidney allografts, such that graft survival in the first year after transplantation now exceeds 90% [1]. However, improvement rates have decelerated. Data from 135 kidney transplant centers in 21 European countries (187,787 individual transplantations) indicate that the improvement of graft survival has slowed significantly since 2000, even when considering the increased age of donors and recipients [1]. Initiatives to further improve graft survival rates are therefore needed, but the nature of such initiatives is an important point for discussion.
Certainly, several pharmaceutical regimens have been developed, such as efficacious and relatively well-tolerated immunosuppressants, which have enabled very good short-term outcomes to be achieved in patients (and with an acceptable risk of graft rejection). However, the ensuing misconception is that all major hurdles in transplantation have been overcome [2, 3]. Good short-term outcomes that are observed in transplantation recipients do not always translate into satisfactory long-term graft functioning or patient-survival rates [4, 5]. Lack of long-term success creates difficulty in defining suitable surrogate endpoints for clinical trials, which is problematic for ongoing research and could discourage academic/commercial investment in kidney transplantation [3].

Consequently, while clinical progress in kidney transplantation is slowing down, rates of allograft loss continue to be unacceptably high. The extensive negative impact that this has on patient health and well-being—as well as the high long-term health-associated cost burden [6]—clearly indicates a need for novel, effective management strategies, tested according to endpoints that suit current practice and regulations. In addition, there are no approved surrogate markers for long-term graft failure in kidney transplantation, necessitating long-term interventional studies as an urgent priority.

Innovations in kidney transplantation often focus on the prevention and treatment of acute allograft rejection [7]. While current immunosuppressive regimens have reduced the incidence of rejection in low-risk organ recipients [8], many patients have a high immunological risk and could benefit from better preventive and therapeutic options than those available [9]. Stratification of patients and allografts according to immunological risk, however, is not standardized.

**CURRENT EMA GUIDANCE ON CLINICAL STUDIES FOR KIDNEY TRANSPLANTATION**

Released in 2008, the European Medicines Agency (EMA) guideline (CHMP/EWP/263148/06) [10] provides guidance on the conduct of clinical studies for solid organ transplantation (not specific for kidney transplantation) by defining treatment goals, study designs, outcome measures, and data analyses for new immunosuppressive therapies developed to prevent and treat allograft rejection. The guideline [10] defines the primary efficacy endpoint for novel immunosuppressants in solid organ transplantation.

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**BOX 1 | Individuals involved with the EMA-CHMP request for Broad Scientific Advice project, on behalf of the European Society for Organ Transplantation.**

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transplantation (not specific for kidney transplantation) as a composite of four outcomes:

- Patient death
- Graft failure—defined by discrete criteria (e.g., permanent return to pre-transplantation treatment modality for a specific period)
- Biopsy-proven acute rejection—including pathological grading for the transplant, outcome, treatment, and response
- Graft (dys)function—defined by best available clear-cut and discrete criteria for kidney, lung, and heart transplantations (e.g., measurement of creatinine/inulin clearance for kidney dysfunction).

Components of the composite endpoint could be omitted for several reasons, such as if there is limited sensitivity/specificity for available biomarkers, or limited consensus about the importance of individual risk factors or cut-off values. Factors such as previous early graft loss (because of immunological factors), re-transplantation, human leukocyte antigen (HLA) mismatch, and presence of HLA antibodies are often taken into consideration, depending on the type of transplantation. The guideline also notes that best attempts should be undertaken to define the recipient’s immunological risk at baseline, using categories such as “low/medium/high” or “elevated/non-elevated.” Finally, CHMP/EWP/263148/06 notes that transplantation outcome is also influenced by surgery and comorbidity [10]. Therefore, reasonably validated scales for assessment of global transplantation risk are important and should be reflected in the target population of clinical studies.

**RATIONALE FOR UPDATING THE GUIDELINES**

Not only has clinical organ transplantation changed markedly since the publication of CHMP/EWP/263148/06 [10]; the transplant community now observes signs of substantial decline in the rate of clinical innovation in this field. Consequently, ESOT—the umbrella organization under which all European transplant activities are organized—sought to understand whether updating guidelines on clinical trial endpoints might help to encourage kidney transplantation research, since potentially outdated or unclear definitions of risk groups and markers of success or failure might limit investment or innovation in transplantation medicine.

**METHODS**

The project that ultimately resulted in the Broad Scientific Advice request and the present special issue began in May 2016, when the European Medicines Agency responded positively to ESOT’s request to begin interactions relating to the overall topic. Within ESOT, the project was coordinated from its inception by Professor Maarten Naesens of KU Leuven, Belgium, with contributions by an international group of experts. This panel of volunteers included nephrologists, surgeons, transplant pathologists, epidemiologists, immunologists, and researchers (Box 1). The key events in the development process were:

- September 26, 2016: First workshop meeting at ESOT Barcelona to outline the project, define the core questions, and appoint working group leaders
- 2017–2018: Briefing package for the EMA was prepared by attendees of the workshop in Barcelona, who collaborated via telephone/e-mail and personal interaction
- June 11, 2018: Submission of a briefing package to EMA, outlining the scope and core questions put forward by ESOT
- June 20, 2018: Response from EMA and invitation to submit a formal request for Broad Scientific Advice
- 2018–2019: Establishment of five working groups related to the core questions agreed with EMA (Box 2), on which ESOT sought advice
- 2018–2019: Writing of the first drafts of the answers to the questions and searching consensus within the working groups
  - Each working group approached the Centre for Evidence in Transplantation (CET) with specific data extraction requests
  - Information provided by CET was used by each working group at their own discretion to produce draft documents
  - The documents were drafted by the experts within each working group
- September 17, 2019: Workshop at the ESOT congress in Copenhagen: consensus meeting to discuss the conclusions reached by the working groups
- 2019–2020: Finalization of the consensus document and preparation of the official request for Broad Scientific Advice

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**BOX 2 | Questions presented to EMA by ESOT in their request for Broad Scientific Advice on clinical trial design and endpoints in kidney transplantation. This special issue presents current knowledge and perspectives from the European Society for Organ Transplantation (ESOT) on these five core questions, based on evidence and clinical and research experience.**

Q1: Does CHMP agree with the updated definitions of rejection and their potential use as primary endpoints in studies of kidney transplantation?
Q2: Does the CHMP agree with the proposed definitions of allograft (dys)function in kidney transplantation, and the recommendations for parameters that could be used as primary endpoints in clinical trial settings?
Q3: Does CHMP agree with the proposed specific risk profiles for kidney transplantation which determine background risk of rejection associated with immunosuppressive therapy?
Q4: Does CHMP agree that long-term outcome after kidney transplantation is an area of unmet medical need, for which conditional marketing authorization procedures should be considered, to facilitate timely access to new therapies? If so, does CHMP agree with the proposed surrogate endpoints for clinical trials for therapies requiring conditional marketing authorization?
Q5: Does CHMP agree with the proposed patient reported outcomes as (primary/secondary) endpoints for use in clinical trials of kidney transplantation interventions?
June 5, 2020: Submission of the package to request Broad Scientific Advice from CHMP
June 6–11, 2020: Start of the Scientific Advice Working Party (SAWP) procedure
July 6–9, 2020: SAWP discussion meeting, at which a list of issues to be addressed by ESOT was adopted
September 24, 2020: ESOT submitted additional documentation to the SAWP
September 30, 2020: Virtual discussion meeting between ESOT and the SAWP, addressing the list of issues
September 28 to October 1, 2020: SAWP agreed on the advice to be given to ESOT
December 7–10, 2020: adoption of the advice by CHMP and response received by EMA
2020–2021: Reformattting of the package submitted to CHMP to fit publication style (in article form and as a positioning paper), to allow widespread dissemination of ESOT’s answers to the questions and the CHMP advice
2021: The ESOT position on each question, including any comments from EMA, is summarized in evidence-based articles within this special issue.

CONCLUSION

In June 2020, ESOT presented an extended form of the articles and supporting materials in this special issue to the EMA as one document, as part of the package to request Broad Scientific Advice from CHMP. Following constructive responses from the EMA, this special issue is published to communicate outcomes and extend discussions among the wider kidney transplant community. Ultimately, it is hoped that endpoints suitable for future clinical trials and better consensus on risk stratification are developed and agreed globally, thereby increasing the quality of future research and evidence, and advancing the practical management of kidney transplantation, particularly for long-term outcomes.

AUTHOR CONTRIBUTIONS

This article is one of a series of papers developed from content relating to the Broad Scientific Advice request, submitted to the European Medicines Agency (EMA) Committee for Medicinal Products for Human Use (CHMP) by the European Society for Organ Transplantation (ESOT) in 2020: interactions between the EMA and ESOT regarding this request began in 2016. The present article was adapted by MN from the final Broad Scientific Advice request submission (June 2020), presentation documents and minutes of the meeting between ESOT and the CHMP Scientific Advice Working Party (SAWP) (September 2020), and the final response from the SAWP (December 2020), to form an introduction to the special issue and present the rationale and methodology used throughout the process; the article was reviewed by SS and finalized by both co-authors.

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CONFLICT OF INTEREST

The authors declare that the work was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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