Topic: Emerging translational safety technologies and tools for interrogating human immuno-biology

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Topic details

Action type: Research and Innovation Action (RIA)
Submission and evaluation process: 2 stages

Specific challenges to be addressed

There is an urgent need to better understand inherent risks of innovative therapeutics for immuno-oncology and immuno-inflammatory disease indications including cytokine release syndrome, infection, malignancy and autoimmunity during early (non-clinical) stages of drug development. The toxicities induced by immunomodulatory therapeutics in patients are often not detected in the young healthy animal models that are routinely used for toxicology studies either due the lack of expression of appropriate drug targets/pathways or due to differences in drug target genetics, expression and functions between animal models and the intended patient populations. Thus, innovative translational safety assessment tools, technologies, models and resources are needed to facilitate the development of novel immunomodulatory drugs (either immunostimulatory or immunosuppressive). Improving the predictivity of non-clinical models will help deliver safer efficacious immunomodulatory medicines to patients and contribute to the principles of the 3Rs (i.e replacement, reduction, and refinement of the use of animals for research). The topic requires a strategic consortium that will enable the sharing of experience from regulators, academia, and pharma industry, in fast evolving immune cell phenotyping technologies, complex in vitro model development, and characterisation of engineered and disease state animal models, as well as facilitating access to extensive immune cell and tissue biobanks (including patient-derived material). Immunomodulatory therapeutic modes of action (spanning both oncology and non-oncology therapeutic indications) for which there is existing non-clinical and clinical safety experience within the pharma industry, academia and health authorities, provide a powerful framework within which the utility of innovative non-clinical models and endpoints can be assessed for potential use in future Investigational New Drug (IND)/Clinical Trial Application (CTA)-enabling safety assessment packages.

Need and opportunity for public-private collaborative research

This topic focuses on a defined subset of immunomodulatory therapeutic modes of action (spanning both oncology and non-oncology therapeutic indications) for which there is existing non-clinical and clinical safety experience within the pharma industry, academia and health authorities. The assessment of the potential utility of integrating safety data derived from innovative non-clinical animal models and/or human in vitro immune cellular system into IND/CTA is unlikely to be within the scope of investment and capabilities for any one company or organisation, but collaborative efforts in a public-private partnership are most likely to harness all the skills and expertise required. Such non-clinical models and endpoints will need to be customised for specific immunomodulatory therapeutic modes of action, disease indications and/or anticipated toxicities. The importance of early engagement and alignment plans with a broad range of public and private stakeholders is exemplified by the themes discussed in several recent workshops and publications on the safety assessment of innovative immunomodulatory therapeutics [1][2][3].
Scope

This proposal aims to establish a public-private consortium that will enhance translational safety assessment approaches for immunomodulatory therapeutics (spanning oncology and non-oncology indications) through development and validation of innovative non-clinical tools and technologies, supported by access to in vitro, ex vivo and in vivo model-derived immune cell and tissue resources, with an emphasis on evaluating human-relevance. Potential toxicities induced by immunomodulatory therapeutics in patients are often not evident in traditional animal models due to lack of expression of appropriate targets, pathways and/or differential expression/functions versus intended patient populations (e.g. species differences in immune cell repertoire and functions including checkpoints and MHC/peptide complex formation). Patient-derived and advanced animal models should thus enable a better understanding of the interplay between drug target and disease state-associated immune cell repertoires.

At present, in vitro human immune cell assays currently used to assess potential effects of immunomodulatory therapeutic agents on functional endpoints such as cytokine release or T cell activation are typically derived from healthy animals and human donors. Thus, there is a need to more accurately model the complex signaling interactions between multiple immune cells in both healthy and disease states (e.g. tumor microenvironment). Therefore the generation of comparative (cross-species) molecular, biochemical, cellular immunobio-logic resources coupled to functional and phenotypic outcomes should provide better definition of limitations in the translatable of in vivo and in vitro test systems to patients and guide optimal non-clinical safety assessment strategies based on therapeutic target, modality, disease indication(s) and patient susceptibility factors (e.g. genotype; biomarkers of immune memory).

Moreover, investing in novel human relevant immune-competent microphysiological systems to assess risks associated with immunomodulatory drugs (e.g. immune-related adverse events, infection, malignancy) will contribute to the effort of reducing animal use while improving predictivity of preclinical models.

The following objectives are within the scope of this proposal:

1. development of innovative comparative (cross-species) in situ and ex vivo molecular, biochemical tools and cellular profiling of immune cells (including patient-derived clinical samples) and association with functional/phenotypic endpoints to enable:
   - enhanced understanding of therapeutic target and pathway biology,
   - enhanced pharmacological and toxicological mechanistic insight and biomarker identification supporting therapeutic index determination for drug candidates and drug modalities,
   - identification of novel endpoints/biomarkers to help support safe starting dose selection for first in human studies e.g. determination of minimum pharmacologically active dose (mPAD) and minimum anticipated biological effect level (MABEL);

2. establishment, refinement and validation of non-clinical tools and models to enable the development of novel classes of immunomodulatory medicines supporting in vitro-in vivo and cross-species translation:
   - comparative assessment of performance of established human in vitro immune cell assays (e.g. cytokine release assays; T cell activation and target cell interaction assays) based on healthy volunteer versus patient-derived cells representing specific disease states,
   - development and validation of human immune-competent organotypic and microphysiological systems for efficacy and safety profiling of immunomodulatory drugs (including back translation of clinical findings),
   - development and validation of animal models (e.g. humanised mice, genetically engineered animal models, disease models) as tools for efficacy and safety assessment of immunomodulatory drugs (including refined models to predict risk of infection / malignancy / immune-related adverse events),
deployment of computational biology approaches for assessment and integration of large multivariate immuno-biology datasets (e.g. phenotypic-anchoring of molecular, biochemical and cellular endpoints derived from both non-clinical models and clinical trials) to reveal novel immune-biology mechanisms and translational biomarkers.

Given the significant technical and logistical challenges associated with sourcing patient-derived samples to support customised safety assessment studies, the topic will also aim to establish a network to facilitate sourcing of human-derived immune cell and tissue samples via centres of excellence for specific disease areas (patient organisations / medtech / academic partners).

Expected key deliverables

- Prioritisation of immunomodulatory therapeutic modes of action (MoA) (including immuno-oncology and immuno-inflammatory disease indications) for which there is extensive non-clinical and clinical safety experience but suboptimal prediction of human toxicities based on input from regulators.
- Development/evaluation of innovative molecular and cellular immunophenotyping biomarkers across conventional toxicology models, ‘engineered’ animal models, and human in vitro models. Provide enhanced molecular and cellular biomarkers for determination of therapeutic index (TI) and first-in-human (FIH) maximum recommended starting dose (MRSD). Benchmark biomarker performance versus patient-derived samples and assess utility for determination of minimum pharmacologically active dose (mPAD) and minimum anticipated biological effect level (MABEL).
- Development/evaluation of human in vitro systems (healthy donor- and/or patient-derived) and ‘engineered’ animal models that are customised for specific therapeutic MoA and patient-relevant cell types/tissue microenvironments in order to recapitulate target engagement, pharmacodynamics and clinically-relevant toxicity phenotypes.
- Leverage consortium approach to enable sharing of experience, tools, models, biobanked samples and databases between regulators, academia and pharma industry (contributing to the principles of the 3Rs).
- Establishment of a network to facilitate sourcing of human-derived immune cell and tissue samples for specific disease areas based on input from patient organisations, medical technology companies and academic centres of excellence.
- Development of customised non-clinical safety assessment strategies (e.g. human in vitro / ex vivo models and/or engineered animal models) for immunomodulatory therapeutics based on therapeutic target, modality, disease indication(s), and clinical safety experience; aligned with regulatory expectations.

Expected impact

The action generated by this topic will ultimately help deliver safer medicines to patients via:

- provision of new tools and models to enable a better understanding of the inherent safety risks of immunomodulatory therapeutics during early (non-clinical) stages of drug development;
- improvement of drug development processes and regulatory assessments for immunomodulatory therapeutics through the characterisation of innovative immune-biology models and biomarkers that complement and enhance existing non-clinical safety assessment approaches;
- better definition of limitations in the translatability of non-clinical test systems to patients, which will enable the most appropriate and efficient combinations of test systems to be used for future safety assessment of immunomodulatory therapeutics, and will also contribute to the principles of the 3Rs.

Small and medium-sized enterprises (SMEs) can be of great benefit to IMI2 JU projects. Their involvement in the action might offer a complementary perspective to industry and academia, and help deliver the long-term impact of the project. Therefore, applicants should indicate how their
proposals will impact and strengthen the competitiveness and industrial leadership of Europe by, for example, engaging suitable SMEs.

**Potential synergies with existing consortia**

Applicants should take into consideration, while preparing their short proposal, relevant national, European (both research projects as well as research infrastructure initiatives), and non-European initiatives. Synergies and complementarities should be considered in order to incorporate past achievements, available data and lessons learned where possible, thus avoiding unnecessary overlap and duplication of efforts.

In particular, the action generated by this topic should, among others, consider initiatives such as:

- Collaborative Network For Immunological Safety Research in Minipigs (CONFIRM) Initiative ([https://minipigs.dk/knowledge-base/the-confirm-initiative/](https://minipigs.dk/knowledge-base/the-confirm-initiative/))
- BioSafe Expert Working Group on improving translational value of *in vitro* and animal models for assessing the pharmacology and toxicology of ImmunoOncology therapeutics ([https://www.bio.org/events/biosafe-meeting-agenda](https://www.bio.org/events/biosafe-meeting-agenda))
- MIROCALS: Efficacy and safety of low-dose IL-2 (ld-IL-2) as a Treg enhancer for anti-neuroinflammatory therapy in newly diagnosed Amyotrophic Lateral Sclerosis (ALS) patients ([http://www.mirocals.eu/](http://www.mirocals.eu/))

**Industry consortium**

The industry consortium is composed of the following EFPIA partners:

- Novartis (lead)
- Roche (co-lead)
- Bluebirdbio
- Boehringer-Ingelheim
- Covance
- MerckSerono
- Sanofi
- Servier
- Transgene AG

In addition, the industry consortium includes the following IMI2 JU Associated Partner:

- JDRF

The industry consortium will include expertise in *in vitro* and *in vivo* safety assessment models and biomarkers and will contribute mainly in the form of:
design and deployment of prospective in vitro and in vivo studies; provision of historical safety-related in vivo phenotypic data for defined therapeutic modes of action (e.g. discontinued development compounds; marketed compounds); in silico prediction of immunomodulatory drug-related adverse events;

provision of advanced technology platforms and bioinformatic support for cross-species molecular, biochemical and cellular phenotyping of immune cells e.g. spatial molecular profiling of RNA/proteins/metabolites; integrated transcriptomic and epigenomic profiling of immune cell subsets enriched by flow cytometry and/or at single cell resolution; mass cytometry characterisation of signalling pathways;

provision of human blood- and tissue- samples to support deployment of advanced immune cell phenotyping technologies and/or in vitro model development (baseline, antigen-challenged, pharmacologically-stimulated, disease states);

emerging human-centric ex-vivo / in vitro technologies (patient-derived and healthy volunteer-derived) will be evaluated and/or further refined for their potential to recapitulate in vivo human immune biology (e.g. in vitro modelling of human immune responses including impact of genetic variants and immunogenicity; 3-D bioprinting and microfluidic technologies, including organ-on-a-chip or hollow-fiber systems and spheroid/organoid modelling, to enable derivation of complex organotypic and microphysiologic systems.

Indicative duration of the action

The indicative duration of the action is 72 months.

Indicative budget

The indicative industry in-kind contribution is EUR 11 000 000.

This contribution comprises an indicative EFPIA in-kind contribution of EUR 10 895 000 and an indicative IMI2 JU Associated Partners in-kind contribution of EUR 105 000.

Due to the global nature of the participating industry partners and IMI2 JU Associated Partners, it is anticipated that some elements of the contributions will be non-EU/H2020 Associated Countries in-kind contributions.

The financial contribution from IMI2 JU is a maximum of EUR 11 000 000.

Applicant consortium

The applicant consortium will be selected on the basis of the submitted short proposals. The applicant consortium is expected to address all the objectives and make key contributions to the defined deliverables in synergy with the industry consortium which will join the selected applicant consortium in preparation of the full proposal for stage 2. This may require mobilising, as appropriate the following expertise and resources:

regulatory expertise (either as project beneficiary or member of a Regulatory Advisory Board of the project) on selection of immunomodulatory therapeutic case studies for assay/model validation and for refining criteria used to make future project decisions e.g. validation using existing examples of discontinued compounds based on clinical evidence/outcomes. Regulators from global health authorities will contribute to the selection of representative immunomodulatory therapeutic modes of action for which there is perceived to be an opportunity to enhance translational safety assessment;

expertise in customised/innovative immune cell assay/model/bioinformatics development;

expertise in innovative technology platforms for molecular, bioinformatics development of immune cells;

provision of human blood- and tissue- samples to support deployment of advanced immune cell phenotyping technologies and/or in vitro model development;
ability to develop ex-vivo / in vitro technologies for recapitulating in vivo human immune-biology;

- given the significant technical and logistical challenges associated with sourcing patient-derived samples to support customised safety assessment studies, significant experience in the establishment of a network to facilitate sourcing of human-derived immune cell and tissue samples via academic and clinical centres of excellence for specific disease areas that may include engagement of relevant patient organisations.

In addition to academic groups, relevant SMEs with relevant proven expertise are encouraged to participate in the applicant consortium. SMEs can be of great benefit to IMI projects and, inter-alia strengthen the competitiveness and industrial leadership of Europe. Their involvement might offer a complementary perspective to industry and academia, and strengthen the long-term impact of the project. For these reasons, applicants should consider engaging SMEs throughout the proposal. Under this topic, the contribution of SMEs would be considered especially beneficial in the provision of innovative, engineered animal models and/or in vitro models that mimic human immuno-biology.

The size of the consortium should be proportionate to the objectives of the topic.

Suggested architecture of the full proposal

The applicant consortium should submit a short proposal which includes their suggestions for creating a full proposal architecture, taking into consideration the industry contributions and expertise provided below.

The final architecture of the full proposal will be defined by the participants in compliance with the IMI2 JU rules and with a view to the achievement of the project objectives.

In the spirit of the partnership, and to reflect how IMI2 JU call topics are built on identified scientific priorities agreed together with EFPIA beneficiaries/large industrial beneficiaries, these beneficiaries intend to significantly contribute to the programme and project leadership as well as project financial management. The final architecture of the full proposal will be defined by the participants in compliance with the IMI2 JU rules and with a view to the achievement of the project objectives. The allocation of a leading role within the consortium will be discussed in the course of the drafting of the full proposal to be submitted at stage 2. To facilitate the formation of the final consortium, until the roles are formally appointed through the consortium agreement, the proposed project leader from among EFPIA beneficiaries/large industrial beneficiaries shall facilitate an efficient negotiation of project content and required agreements. All beneficiaries are encouraged to discuss the project architecture and governance and the weighting of responsibilities and priorities therein.

The consortium is expected to have a strategy on the translation of the relevant project outputs into regulatory, clinical and healthcare practice. A plan for interactions with regulatory agencies / health technology assessment bodies with relevant milestones and resources allocated should be proposed to ensure this (e.g. qualification advice on the proposed methods for novel methodologies for drug development, qualification opinion).

A plan for aspects related to sustainability, facilitating continuation beyond the duration of the project should also be proposed.

The architecture outlined below for the full proposal is a suggestion. Different innovative project designs are welcome, if properly justified.

Work package 1 – Management, coordination, dissemination and sustainability

The goals of this work package will be as follows:

- overall coordination;
- liaise with regulatory advisors to establish a framework for selection of immunomodulatory therapeutic modes of action that will be assessed using new models and biomarkers during the project. The selection of specific therapeutic modes of action / drugs for evaluation in innovative
non-clinical models will be managed via a face-to-face full consortium meeting (including regulatory advisory board input) during the first few months of the active project timeline (i.e. post-launch);

- dissemination of scientific results and research data;
- active communication of promising biomarkers and/or non-clinical models for the safety assessment of immunomodulatory therapeutics into the public domain via stakeholder workshops, publications and presentations;
- development of a sustainability plan.

**Expected applicant consortium contribution:** project management including coordination of work package deliverables, periodic reporting and budget administration, dissemination of scientific results and development of a sustainability plan.

**EFPIA consortium contribution:** leadership of overall project goals with respect to safety assessment applications, communication, dissemination of project results and development of sustainability plan.

**Work package 2 – Innovative models for safety assessment of immuno-oncology therapeutics**

The goals of this work package will be as follows:

- selection of clinically validated immunomodulatory therapeutic modes of action (including both small molecule and biotherapeutic immuno-oncology therapeutics) for which there is perceived to be an opportunity to enhance translational safety assessment;
- deployment of customised non-clinical models (with a particular emphasis on improved modelling of contributions from tumour target antigens and tumour microenvironments including biological barriers such as the intestine epithelium or the blood-brain barrier) to complement existing clinical and non-clinical safety profiles. It is envisaged that a customised combination of animal models and human in vitro models will need to be deployed for enhanced characterisation and/or prediction of clinical toxicities associated with specific immunodulatory therapeutic modes of actions / drug targets.

**Expected applicant consortium contribution:** co-lead work package, study design, provision/deployment of models and/or biomarkers, and dissemination of results.

**EFPIA consortium contribution:** co-lead work package, study design, provision/deployment of models and/or biomarkers, and dissemination of results.

**Work package 3 – Innovative models for safety assessment of immuno-inflammatory disease therapeutics**

The goals of this work package will be as follows:

- selection of clinically validated immunomodulatory therapeutic modes of action (including small molecule and biotherapeutic immuno-inflammatory disease oncology therapeutics) for which there is a perceived opportunity to enhance translational safety assessment;
- deployment of customised non-clinical models (with a particular emphasis on improved modelling of contributions from inflammatory disease target biology and tissue microenvironments including biological barriers such as the intestine epithelium or the blood-brain barrier) to complement existing clinical and non-clinical safety profiles. It is envisaged that a customised combination of animal models and human in vitro models will need to be deployed for enhanced characterisation and/or prediction of clinical toxicities associated with specific immunodulatory therapeutic modes of actions / drug targets.

**Expected applicant consortium contribution:** co-lead work package, study design, provision/deployment of models and/or biomarkers, and dissemination of results.
EFPIA consortium contribution: co-lead work package, study design, provision/deployment of models and/or biomarkers, and dissemination of results.

**Work package 4 – Innovative biomarkers for translational safety assessment of Immunomodulatory therapeutics**

The goals of this work package will be as follows:

- develop innovative comparative (cross-species) *in situ* and *ex vivo* molecular, biochemical tool and cellular profiling of immune cells (including patient-derived clinical samples) and association with functional/phenotypic endpoints (e.g. autoantibodies, cytokine release, skin toxicity, neurotoxicity);
- deploy computational biology approaches for assessment and integration of large multivariate immuno-biology datasets (e.g. phenotypic-anchoring of molecular, biochemical and cellular endpoints derived from both non-clinical models and clinical trials) to reveal novel immune-biology mechanisms and translational biomarkers;
- apply established discovery and validation approaches for biomarker development in order to ultimately contribute to facilitating patient stratification and monitoring based on biopsy and minimal invasive liquid biopsy testing.

Expected applicant consortium contribution: co-lead work package, study design, provision/deployment of biomarker analytical and analysis technologies, and dissemination of results.

EFPIA consortium contribution: co-lead work package, study design, provision/deployment of biomarker analytical and analysis technologies, and dissemination of results.

**Work package 5 – Clinical sample management – sourcing and logistics for assay and model development**

The goals of this work package will be as follows:

- establish an IMI project-coordinated network to facilitate sourcing of human-derived immune cell and tissue samples via centres of excellence for specific disease areas;
- establish appropriate logistics for sample tracking and biobanking;
- establish a framework for ensuring appropriate ethical and legal review of proposed consortium research use of human-derived cells and tissues;
- qualify pre-analytical procedures for sample collection and implementation of novel workflows/devices, minimising pre-analytical variances for enabling reliable molecular diagnostics, implement/confirm multi-modal & multi-analyte diagnostic approaches.

Expected applicant consortium contribution: Lead work package, engage key stakeholders from academic clinical centres of excellence and patient organisations, establish an ethical review board.

EFPIA consortium contribution: Co-lead for work package, definition of specific human immune cell and tissue samples that would support model development, provision of guidance for sample tracking and biobanking.

**References**
