Optimising *in vitro* methods through removal of serum components

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Increasing acceptance of the role *in vitro* • assays can play in assuring consumer safety, particularly as part of Next **Generation Risk Assessment (NGRA)**

Growing desire to remove animal products lacksquarefrom these in vitro assays to make them more scientifically robust and humanrelevant.



Safety & Environmental Assurance Centre



Towards the Development of Animal Product-Free in vitro Systems for NGRA of **Consumer Goods** Paul Russell, Sarah Hatherell, Ouarda Saib & John Kilgallon

Background

There is an increasing acceptance of the role in vitro assays can play in assuring consumer safety, particularly as part of Next Generation Risk Assessment (NGRA) (Baltazar et al, 2020), NGRA is an exposure-led, hypothesis-driven approach integrating new approach methodologies (NAMs) to ensure chemical safety without the use of animal testing. There is also a growing desire to remove animal products from these in vitro assays to make them more scientifically robust and human relevant. For example, the use of foetal bovine serum (FBS) and animal-derived, antibodies can introduce a lot of batch-to-batch variability potentially resulting in experimental quality (e.g. contamination of FBS; specificity of antibodies) and reproducibility issues (Baker et al, 2016; van de Valk et al, 2018). Additionally, it is more frequently becoming recognised that knowledge of all the constituents of the cell culture medium used and their influence on cellular processes are important for improved reproducibility (Hirsch & Schildknecht, 2019). Therefore, ideally chemically defined media would be used to culture human cells for in vitro assays to eliminate any remaining scientific quality issues resulting from use of animal- or human-derived components (van der Valk et al, 2010) although this is technically very challenging. Here we will describe some of the challenges, opportunities and potential options for replacing animal-derived products in in vitro systems.

Key Focus Areas for Replacement of animal-derived products

Cell Culture Media/Reagents

In 2008, the ECVAM Scientific Advisory Committee (ESAC) advocated for the use of non-animal derived supplements for in vitro studies wherever possible and stated that "for methods forwarded to ECVAM for validation/pre-validation where [the use of non-animal alternatives to serum] is not fulfilled a justification for future use must be provided, including measures taken to seek non-animal alternatives to [FBS]

The technical disadvantages of using serum (both animal & human-derived) include its undefined nature, batch-to-batch variability in composition, and the risk of contamination. Several OECD test guidelines* already use serum-free cell culture methods demonstrating that this is an area of growth with plenty of opportunities for further development. In particular it is hoped that rather than just removing or replacing (i.e. animal with human) serum in cell culture medium a fully chemically defined medium can be used as there are also reproducibility concerns due to the non defined nature of human serum

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Current challenges include commercial availability of specialised reagents; technical challenges when it comes to cell adaptation (each cell type requires its own specific media) which requires expertise and can be a lengthy process; ensuring that the whole protocol is animal product free (i.e. from the plastic ware to the reezing media); lack of consensus around what constitutes an animal-free reagent

ies include the growing market for those who want more reproducible, reliable, ethical and human-relevant assays; faster research; improved assays, *including QECD TGs no. 431, 439 and 492 (QECD, 2015a: QECD, 2015b: QECD, 2016) in vitro test methods for skin corrosion, skin irritation & eve irritation testing

Antibodies

The recent EURL ECVAM paper on animal derived antibodies (Viegas Barosso et al. 2020) sets out a clear position: "EURL ECVAM recommends that animals should no longer be used for the development and production of antibodies for research, regulatory, diagnostic and therapeutic applications. In the EU, the provisions of Directive 2010/63/EU should be respected, and EU countries should no longer authorise the development and production of antibodies through animal immunisation, where robust, legitimate scientific justification is lacking.

A clear scientific advantage of using non-animal derived antibodies is again the removal of batch-to-batch variability. The main challenge here is that animal derived antibodies are currently much more readily available as "off the shelf" and ready to use than those that don't involve animals

We recognise that transition to routine use of non-animal-derived antibodies will be greatly helped by the availability of mature technologies and specialised laboratories/manufacturers. The hope is to work with those developers and other external collaborators to increase the uptake and use of these types of antibody, driving further development

Existing collaborations

Unilever is sponsoring (with AstraZeneca) and co-funding NC3Rs CRACK IT challenge 36 ps://nc3rs.org.uk/crackit/animal-free-vitro) to adapt two established OECD Test Guideline in vitro systems to animal-product-free conditions.

The two in vitro tests are OECD TG 487 and OECD TG 455: they are important tests as they provide data towards establishing whether a chemical will cause cancer or endocrine disrupting effects in humans

The aim of this work is to deliver a robust, human-relevant (and preferably chemicallydefined) version of the assays that demonstrates improved data quality and reproducibility.

further so that the tests are performed under fully chemicallydefined conditions **XCellR8** Redefining testing

sensitisation in vitro tests KeratinoSens™ (OECD TG442D) and

We are currently collaborating with XCellR8 to take this a step

with human serum (Belot et al. 2017: Edwards et al. 2018)

h-CLAT (OECD TG442E) to xeno-free conditions by replacing FBS

XCellR8 have previously successfully adapted two skir

Conclusion: Increased development, publication and commercial availability of animal-free reagents will increase the reproducibility and human relevance of the chosen assays and help drive the use of animal product-free in vitro systems for NGRA.

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Technical disadvantages of using serum...

...include its undefined nature, batch-to-batch variability, & risk of contamination. (S167 – Replacing Fetal Bovine Serum)

In particular it is hoped that rather than just removing or replacing (i.e. animal with human) serum in cell culture medium a **fully chemically defined medium** can be used.

Chemically defined media

Standard *in vitro* cell culture media commonly consist of a basal medium supplemented with serum (animal or human-derived) as a source of nutrients and other ill-defined factors. In contrast, chemically defined media require that **all of the components must be identified and have their exact concentrations known**.



Non-Animal-Derived Antibodies (NADAs)

"EURL ECVAM recommends that animals should no longer be used for the development and production of antibodies for research, regulatory, diagnostic and therapeutic applications."

Viegas Barroso et al, 2020 EURL ECVAM Recommendation on Non-Animal-Derived Antibodies.



A clear **scientific advantage** of using NADAs is again the **removal of batch-to-batch variability**.

Main **challenge** here is that animal-derived antibodies are currently much more readily available as "off the shelf" and ready to use than NADAs.

Recognise that transition to routine use of NADAs will be greatly helped by the **availability of mature technologies and specialised laboratories/manufacturers** (S118 – A global movement to improve science using animal-free antibodies)

Hope to work with developers & other external collaborators to **increase the uptake & use** of these types of antibody, **driving further development**.



Animal Product-Free Cell Culture Media/Reagents

Current challenges

- commercial availability of specialised reagents;
- technical challenges when it comes to cell adaptation (each cell type requires its own specific media) which requires expertise & can be a lengthy process;
- ensuring that the whole protocol is animal product-free (i.e. from the plastic ware to the freezing media);
- lack of consensus around what constitutes an animal-free reagent.

Opportunities - growing market for those who want more **reproducible**, **reliable**, **ethical & human-relevant assays**.



Importance of collaboration

Increased **development**, **publication & commercial availability** of animal-free reagents will increase the **reproducibility & human relevance** of the chosen assays and help drive the use of animal product-free *in vitro* systems for NGRA.

Existing collaborations

Unilever is sponsoring (with AstraZeneca) and co-funding NC3Rs CRACK IT challenge 36 (https://nc3rs.org.uk/crackit/animal-free-vitro) to adapt two established OECD Test Guideline *in vitro* systems to animal-product-free conditions.

The two *in vitro* tests are OECD TG 487 and OECD TG 455; they are important tests as they provide data towards establishing whether a chemical will cause cancer or endocrine disrupting effects in humans.

The aim of this work is to deliver a robust, human-relevant (and preferably chemicallydefined) version of the assays that demonstrates improved data quality and reproducibility. XCellR8 have previously successfully adapted two skin sensitisation *in vitro* tests KeratinoSens^{IM} (OECD TG442D) and h-CLAT (OECD TG442E) to xeno-free conditions by replacing FBS with human serum (Belot et al, 2017; Edwards et al, 2018)

We are currently collaborating with XCellR8 to take this a step further so that the tests are performed under fully chemicallydefined conditions.



