Paving the way for a UK Roadmap:
Development, Validation and Regulatory Acceptance of New Approach Methodologies (NAMs) in Chemical Risk Assessment
Executive summary

Advances in biology, computer science and other related fields are paving the way for major improvements in how we evaluate environmental and public health risks posed by potentially toxic chemicals. The combined advances in discovery and clinical sciences, data science and technology have resulted in toxicity testing which has reached a pivotal transformation point known as the 4th industrial revolution (4IR). One of the major recent scientific advancements is the development of new approach methodologies (NAMs) including computer modelling strategies for the evaluation of hazard and exposure whilst championing the Replacement, Reduction and Refinement (3Rs) of animals, approach.

The volume of data produced in the world is growing ever more rapidly, from 33 zettabytes in 2018 to an expected 175 zettabytes in 2025 (IDC, 2018) (Food Systems). The Department for Business, Energy and Industrial Strategy (BEIS) white paper on Regulation for the Fourth Industrial Revolution notes that changes in technology are occurring at a "scale, speed and complexity that is unprecedented". The use of such technologies can help improve regulatory processes in several ways such as to improve the efficiency of data collection and to exploit data already held by agencies to support better analysis and risk assessment (BEIS Report-The use of emerging technologies for regulation).

The future of food safety assessment of chemicals depends on our adaptability and flexibility whilst using the best scientific methodologies and strategies available in order to respond to the accelerating developments in science and technology. The vision is to be able to predict risk more rapidly and efficiently.

For regulatory agencies to incorporate and implement these new predictive capabilities brings both challenges and opportunities. Moving from research to risk assessment to regulatory setting and beyond, there must be suitable validation and acceptance of these new and emerging technologies.

Using a validation process via an evidence driven approach to address the data gaps in the risk assessment process will facilitate the acceptance and validity of these NAMs as well as pave the way for alternative methods testing strategies. Integration of these technologies as part of the risk assessment process will be fundamental in the future of human and environmental safety.

In order to achieve this, the Food Standards Agency (FSA) and Committee on Toxicity of Chemicals in Food, Consumer Products and the Environment (COT) have developed a UK roadmap towards acceptance and integration of these new approach methodologies including predictive toxicology methods using computer modelling into safety and risk assessments for regulatory decision making.
Introduction & Background

Chemical Landscape

“The Toxicity Data Landscape for Chemicals” paper (Judson et al., 2009) reported that 28 million chemicals had been discovered. Only three million had been tested on animals and some of these also tested in human studies. Approximately, a further one million had some in vitro/in silico screening assay data only. By 2020, over 350,000 chemicals and mixtures of chemicals had been registered for production and use worldwide. The identities of many chemicals remain publicly unknown because they are claimed as confidential (over 50,000) or ambiguously described (up to 70,000) (Wang et al., 2020).

It is understandable that with those numbers, finding an accurate and optimized model poses a big challenge, not least for example some of the environmental chemicals that don’t necessarily come with a toxicological package. However, by combining data from traditional methodologies with data from these new emerging technologies we will be able to predict risk more rapidly and efficiently.

Hopefully this will be an opportunity to engage cross cutting themes, continue to collaborate with the National Centre for the Replacement, Refinement and Reduction of Animal Research (NC3Rs) and tap into other projects e.g., Genome UK: the future of healthcare (2020) and Accelerating the Pace of Chemical Risk Assessment (APCRA) as well as collaborations across academia, industry and beyond.

Current Chemical Challenges

- Too many chemicals to test
- Traditional methods can be slow and costly
- Adopting the principles of the 3Rs (i.e. Replacement, Reduction and Refinement of animal experiments)
- Not enough data on compound/compounds being assessed

Opportunities

Adopt new approach methodologies (NAMs) to predict risk more rapidly, accurately, and efficiently.
What are NAMs and why is there a drive in the regulatory context?

Advances in biology, computer science and other related fields are paving the way for major improvements in how we evaluate environmental and public health risks posed by potentially toxic chemicals. The combined advances in discovery and clinical sciences, data science and technology have resulted in toxicity testing which has reached a pivotal transformation point known as the 4th industrial revolution (4IR). One of the major recent scientific advancements is the development of NAMs including high throughput screening, omics and in silico computer modelling strategies (e.g. Artificial Intelligence (AI) and machine learning) for the evaluation of hazard and exposure. This also advocates the Replacement, Reduction and Refinement (3Rs) of animals, approach (Hartung, 2010) proposed 50 years ago by Russell and Burch (1959).

What is being done?

To keep pace with the digital evolution, we are using the latest technology and best available scientific methodologies to incorporate additional tools into our regulatory risk assessment process to evaluate safety in food, consumer products and the environment more efficiently and without compromising quality.

Why?

This will enable us to provide improved risk assessments of chemicals for which there are currently no, or very few data, and therefore increase consumer safety. NAMs will help us to predict risk more rapidly, accurately, and efficiently.

Who are the UK FSA and the COT?

The UK FSA is an independent Government department working across England, Wales, and Northern Ireland to protect public dietary health and consumers’ wider interests in food. The FSA uses expertise and influence so that people can trust that the food they consume is safe and is what it says it is. The Science, Evidence and Research Division (SERD) of the FSA provides strategic analysis, insight, and evidence across the FSA’s remit to underpin the development of policies, guidance, and advice on food safety.

The COT assesses chemicals for their potential to harm human health. Scientific evaluations are carried out at the request of the FSA, Department of Health and Social Care (DHSC), UK Health Security Agency (UKHSA), and other government departments and regulatory authorities.

Why are the FSA and COT taking a lead on this?
The FSA responds to food incidents and it is imperative that risk assessments on the safety of a chemical are provided. The data used can only be what is available and sometimes there is very little, or no, toxicological information for a given chemical.

For these chemicals, the use of NAMs will provide a more indicative level of risk and therefore greater confidence in risk assessments that individual compounds can be assessed. This will be fundamental in risk assessment scenarios where limited to no information is available on the toxicity of a chemical.

The FSA and COT have been reviewing the use of NAMs to scope what methods are out there, in order for the best methodologies to be used in risk assessment and understand how these can be incorporated in a regulatory context (Environmental, health and safety alternative testing strategies: Development of methods for potency estimation TOX/2019/70). There is a need for integration of several methodologies which will form part of the integrated approaches to testing and assessment (IATAs).

FSA Digital Vision

In the FSA Chief Scientist Data Science Report (2017), it stated that “Big data and data science bring relatively new tools and techniques to Government analytics”.

In the Science Council Working Group on Data Usage and Digital Technology Final Report to the FSA it stated in one of the recommendations: “Encourage the development of data capabilities and skills across the FSA staff base therefore, we want to embrace the new technologies and enhance capabilities for our staff.”

What have the FSA / COT done so far?

Figure 1. Diagram of what the FSA and COT have done so far in the NAMs space.

The scoping paper “Environmental, health and safety alternative testing strategies:
Development of methods for potency estimation” (TOX/2019/70) was reviewed by the COT in December 2019. The COT were provided with a concise review of currently available methods, which included databases, different kinds of Quantitative structure Activity Relationships (QSAR) methods, adverse outcome pathways (AOPs), High Throughput Screening (HTS), read-across models, molecular modelling approaches, machine learning, data mining, network analysis tools, and data analysis tools using artificial intelligence (AI).

The full workshop reports will be made available through the COT website once published (currently under reserved business).

In addition, this work will complement the work surrounding a new subgroup called the Synthesis and Integration of Epidemiological and Toxicological Evidence Subgroup (SETE) of the Committee on Toxicity of Chemicals in Food, Consumer Products and the Environment (COT) and Committee on Carcinogenicity (COC) which was set up in 2019. Its aim was to review the approaches for synthesising and integrating epidemiological and toxicological evidence that are used by the COT and COC in chemical risk assessments and to provide a pragmatic guidance and transparent reflection of how the COT and COC review data.

Exploring Dose Response (EDR) Workshop Summary

The UK FSA and the COT held an “Exploring Dose Response” workshop (March 2020) in a multidisciplinary setting which included regulatory agencies, government bodies, academia and industry. The workshop provided a platform from which to address and enable expert discussions on the latest in silico prediction models, new approach methodologies (NAMs), physiologically based pharmacokinetics (PBPK), future methodologies, integrated approaches to testing and assessment (IATA) as well as methodology validation. Through case studies (including plastic particles, polymers, tropane alkaloids, selective androgen receptor modulators), the workshop outlined and explored an approach that is fit for purpose when applied to human health risk assessment in the context of future food safety assessment. Furthermore, possible future research, to establish points of departure (PODs) using non-animal alternative models and to improve the use of exposure metrics in risk assessment, was discussed.

Overall conclusions were as follows:

a) The use of pragmatic guidelines/framework for incorporating these models into risk assessment. Using case studies, such as those outlined in the workshop, towards applicability and gaining confidence in the models.

b) Human biomonitoring data will be key to identify realistic snapshots of exposure scenarios as well as big data which need to be linked to human clinical data.

c) Exposure data and exposure science will be key in developing in silico models in risk assessment and to explore the use of exposomics.

d) There should be transparency throughout the process i.e. consumer facing engagement on NAMs.

e) There should be planning, to take these new methods forward using social sciences research and technical research for integration.
Ultimately, it was collectively agreed that integration of these new technologies as part of our risk assessment methodologies, with a validation process throughout, will be key in the acceptance of the models (by regulatory bodies) and will be fundamental in the future of human and environmental safety (Figure 2).

Figure 2. Concluding workflow for integration of NAMs in regulatory risk assessment from the EDR workshop.

**PBPK for Regulators Workshop Summary**

The UK FSA and the COT held a “PBPK for Regulators” workshop (December 2020) in a multidisciplinary setting with delegates from regulatory agencies, government bodies, academia and industry. The workshop provided a platform to enable expert discussions and presentations on the application of PBPK to human health risk assessment in a regulatory context as well as potential future research.

Main overarching conclusions of the PBPK workshop:

a) PBPK modelling tools are applicable in the explored areas of use, and there is some expertise available for their utilisation.

b) PBPK modelling offers opportunities from which to address questions for compounds that are otherwise not solvable.

c) Widespread acceptance amongst regulatory bodies appears to be limited by lack of available in-house expertise.

d) Familiarisation using real world case studies would help in developing more experts in the field and increasing acceptance.

e) In a regulatory context, establishing fitness for purpose for the use of PBPK models require multi-partite discussion and harmonised guidance.

f) Finally, PBPK modelling is part of the wider “new approach methodologies” for risk assessment.
Regulatory Acceptance of NAMs

NAMs and IATAs are rarely accepted by regulatory bodies, however it is clear that we are now at a pivotal point where integration of such (methods/techniques) will be fundamental in more efficient and rapid risk assessment. The key question is how these approaches can be facilitated in a regulatory setting using the supporting technology available. The use of these methods through various case studies as a ‘proof of principle’ concept is becoming apparent.

Worldwide perspectives on emerging technologies

The focus of the 7th annual Global Summit on Regulatory Science (GSRS17) was Emerging Technologies for Food and Drug Safety and the summit publication stated that “Moving forward toward greater integration of emerging data and novel methodologies for chemicals risk assessment will need continuous efforts on capacity building” (Slikkler et al., 2018).

Furthermore, in the recent EU Farm to Fork strategy and the EU Green Deal Food 2030 Pathways for Action (Food Systems and Data 2020) it states: “to valorise emerging technologies, tools, standards and infrastructure for use in food systems”.

The Royal Society of Chemistry recently published a Drivers and scope for a UK chemicals framework (2021) which recommends under regulation: “Be a world leader in the development of New Approach Methods (NAMs) for safety evaluation without the use of animals and devise new risk assessment frameworks to support decision-making using NAMs”.

The future direction of safety assessment science will depend heavily on the evolution of the regulatory landscape. A key challenge, though, is whether the regulatory framework can keep pace with the increasing speed of scientific and technological developments (Worth et al., 2019).

Food authorities should strive to incorporate the best scientific methods available (Kavlock et al., 2018) into chemical risk assessment. The future of food safety assessment depends on adaptability, flexibility and revolutionary principles in order to respond to the accelerating developments in science and technology.

This implies that close collaboration will be needed between chemists, toxicologists, informaticians, risk assessors and others to develop, maintain and utilise appropriate models. Not only must the different disciplines come together, but also those scientists from industry, academia and regulatory agencies must recognise the commonalities (Cronin et al., 2019). The challenge is to respond to the growing need for adaptable, flexible and even bespoke computational workflows that meet the demands of industry and regulators, by exploiting the emerging methodologies of Tox21 and risk assessment.
UK Government on NAMS, data and emerging technologies

The volume of data produced in the world is growing ever more rapidly, from 33 zettabytes in 2018 to an expected 175 zettabytes in 2025 (IDC, 2018) (Food Systems and Data). The Department for Business, Energy and Industrial Strategy (BEIS) white paper on Regulation for the Fourth Industrial Revolution notes that changes in technology are occurring at a "scale, speed and complexity that is unprecedented". The use of such technologies can help improve regulatory processes in several ways. These include the improvement of the efficiency of data collection and to better exploit data already held by agencies to support improved analysis and risk assessment.

In the ‘Rebuilding a Resilient Britain’ programme one of the departmental Areas of Research Interest (ARIs) marked as a priority was “data science and digital technologies” in the “Changing Systems” theme.

Furthermore, incorporating NAMs into the regulatory context compliments Defra’s research program on systems thinking approach which includes Food systems.

In the “The Integrated Review of Security, Defence, Development and Foreign Policy Global Britain in a competitive age” it states for a collaborate-access framework to guide government activity in priority areas of Science & Technology, such as AI, quantum technologies and engineering biology which has the potential to unlock a step-change in wide ranging applications, which include chemicals.

This UK NAMs roadmap would tie in with the already existing UKRI Non-animal technologies in the UK: a roadmap, strategy and vision (2015) which recommends a need for early engagement with regulators to ensure that non-animal technologies can be used in regulatory risk assessments.
How does the FSA plan to integrate NAMs in the regulatory space?

Figure 3. Landscape to integrate NAMs in the regulatory space

Formulating the problem space

It is well known that designing and structuring your problem space (Goel and Pirolli, 1992) will be fundamental in its outputs (Newell et al., 1993) and success (Goel and Pirolli, 1989). The science of design consists of the use of efficient, available computational techniques (modelling) for finding the best course of action to respond to real situations, or reasonable approximations of real situations (Simon, 1981).

The FSA plans to use problem exploration techniques which includes interactive sessions at FSA-COT-led workshops, to discuss the use of NAMs in risk assessment and define the questions that need to be answered in order to use NAMs in regulatory chemical risk assessment.
Figure 4. Formulating the problem space for project outputs.

Breaking Down the Silos: Drivers and Obstacles

The Malloy et al. (2017) paper recommends supporting trans-sector and transdisciplinary efforts to integrate predictive toxicology “Use existing efforts to bring together regulators, industry, civil society, and academics to agree on testing protocols for nanotechnologies as a model that then could be adopted in other fields”.

Despite advances, the scope and pace of adoption of NAMs, including toxicogenomics tools and data sets in chemical risk assessment, have generally, not met the ambitious expectations of their advocates (Pain et al., 2020).

Regulatory uptake of NAMs has been hesitant and despite notable improvements and new applications, some of the major obstacles remain, such as unconnected silos, interpretation complexities and acceptance of validation by regulators validation.

Incorporation | Adopters of change | Data to Deployment

The incorporation and implementation of these new predictive capabilities by regulatory agencies brings both challenges and opportunities.

Moving from research to risk assessment to regulatory setting and beyond, the validation and acceptance of these new emerging technologies must be ensured. The FSA plan to learn from other regulatory agencies and other conceptual frameworks (e.g. IPBES Conceptual Framework (Diaz et al., 2015) i.e. to structure the syntheses that will inform policy, and to improve comparability across various
assessments carried out at different spatial scales, on different themes, and in different regions) in different settings.

NAMs are gaining traction as a systematic approach (Sturla et al., 2014) to support informed decisions on chemical risk assessment. Adapting how we assess risk will be, and has always been, a challenge, but big changes have, and can, occur through innovation that facilitates and accelerates adaptation (Rogers, 2003). In Attwell’s 1992 paper, it was concluded that a sensible way forward, was to treat a knowledge barrier approach to technology diffusion, as a distinct theory in its own right. The diffusion of technology is reconceptualized in terms of organizational learning, skill development, and knowledge barriers. As knowledge barriers are lowered, diffusion speeds up, and one observes a transition from an early pattern in which the new technology is typically obtained as a service to a later pattern of in-house provision of the technology.

The “rate of adoption is the relative speed with which an innovation is adopted by members of a social system” Rogers (2003). Therefore, throughout the process the FSA will strive to be transparent and engage with the public. Techniques for adoption and integration could include the spherical cow approach in order for maximum understanding for all our stakeholders including the consumer. Data visualisation exploration techniques could be applied (e.g. Mondrian and Manet (Unwin et al., 1996)) to make use of multidimension data. Incorporating data sets such as Open FoodTox into the latest predictive models will increase predictive capacity for chemicals with little or no tox data.

Transition and Integration

Innovative technologies should be reviewed and evaluated once, prior to integration into the risk assessment/as a chemical testing method, as part of the risk assessment strategies for chemical testing for human health and the environment. Using an evidence driven approach, to address the data gaps in the risk assessment process, will facilitate the acceptance and validity of these NAMs, as well as pave the way for alternative testing strategies with confidence.

It is hard for new technologies to be accepted because regulations, infrastructure, user practices and maintenance networks are aligned to the existing technology (Geels, 2002). In order for these new technologies to progress, users have to integrate them into their practices, organisations and routines. This involves learning/training, adjustments and ‘domestication’ (Lie and Sørensen, 1996). It is known that links between technical and social elements provide stability leading to sociotechnical change.

This UK roadmap will pave the way towards acceptance and integration of NAMs into safety and risk assessments from a regulatory perspective and make use of the government R&D Roadmap which will form part of the wider systems thinking approach.
Integration of these technologies as part of the chemical risk assessment process will be fundamental in the future of human safety i.e. the consumer. This will use the best science available to predict and assess risk more rapidly, accurately, and efficiently.
Paving the way for a UK Roadmap: Development, Validation and Regulatory Acceptance of New Approach Methodologies in Chemical Risk Assessment

Stepping and (Mile)Stones - How the FSA got here

Paving the way for a UK Roadmap Journey - How the FSA got here:

1) Output of the Exploring Dose Response Workshop (March 2020)
2) Output of the PBPK Regulators Workshop (December 2020)
3) Output of the discussions with working groups and scientific advisory committees (regular reviews)
4) First outline of the UK Roadmap
5) Output of the Paving the way for the paradigm shift-UK Roadmap Development, Validation and Regulatory Acceptance of New Approach Methodologies workshop (October 2021)
6) Second outline of the UK Roadmap
7) Finalisation of the UK Roadmap
Overall objectives of the roadmap are to:

- identify latest available NAMs for optimal risk assessment
- learn from other regulatory agencies and beyond
  - validate through case studies
- build confidence in NAMs in the regulatory setting
  - develop skills and training
- implement and integrate NAMs in the regulatory setting
The 7 Steps to Integration & Acceptance
The UK Roadmap

**Listening and Learning**

Outputs from workshops organised by the FSA and COT will be discussed with relevant and appropriate committees, working groups and regulatory agencies for opportunity to comment and provide input.

FSA will review what other regulatory agencies and industry have done and what still needs to be done.

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**Identify and Review**

Review the outputs outlining next steps, identifying needs, gaps, and challenges. Identify available NAMs and what the pros and cons for each are and where their strengths lie.

Formulate the problem space.

Identify opportunities and challenges.

Throughout the process continue to review internal and external research and development and how that may impact on the Roadmap.

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**Training**

Customised in-house training and ongoing education through various seminars and conferences.

Skill development: building a resilient organization.

From the perspective of FSA and COT we can then suggest that other regulators / cross government would need similar training.
Research and Development

Support and initiate research to ensure that the most promising technologies are identified, developed, validated, and integrated.

Assess the list of NAMs and other NAMs roadmaps.

Recruitment of Computational Fellow.

Recruitment of Tox AI PhD student.

Collaboration and Dissemination

Cultivate collaborations across sectors and disciplines nationally and internationally. Dissemination of outputs, data sets and models will play an important role in the uptake and acceptance of NAMs. Identifying the acceptance needs through partnerships and establishing a network to support delivery of new predictive toxicology methods.

Network with regulatory agencies and academics in this workspace.

Set up a hub in the FSA and Cross-Government, industry, and academia to disseminate models etc.

Set up a cross Whitehall NAMs working group so we can exchange information in this area.

Review and Recommend

Collate evidence, review, and ensure transparency. Optimizing the evaluation process.

Maintain open transparency throughout.

Citizen science/public consultation on how the public feel on alternative testing strategies.

Present work throughout to working groups and scientific advisory committees.

Regulatory Integration and Acceptance

Advance the integration of emerging predictive toxicology methods and new technologies into regulatory safety and risk assessments.

Integrate the best possible methodologies in risk assessment.

Virtual laboratories acceptance. Integration of NAMs in a regulatory setting.
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# Abbreviations

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<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>3Rs</td>
<td>Replacement, Reduction and Refinement</td>
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<td>4IR</td>
<td>4th industrial revolution</td>
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<td>AI</td>
<td>Artificial Intelligence</td>
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<td>ARIs</td>
<td>Areas of Research Interest</td>
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<td>AOPs</td>
<td>adverse outcome pathways</td>
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<td>COC</td>
<td>Committee on Carcinogenicity</td>
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<td>FSA</td>
<td>Food Standards Agency</td>
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<td>IATAs</td>
<td>integrated approaches to testing and assessment</td>
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<td>HTS</td>
<td>High Throughput Screening</td>
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<td>NAMs</td>
<td>New Approach Methodologies</td>
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<td>NC3Rs</td>
<td>National Centre for the Replacement, Refinement and Reduction of Animal Research</td>
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<td>PBPK</td>
<td>physiologically based pharmacokinetics</td>
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<td>PODs</td>
<td>points of departure</td>
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<td>SETE</td>
<td>Synthesis and Integration of Epidemiological and Toxicological Evidence Subgroup</td>
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<td>SERD</td>
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<td>UKHSA</td>
<td>UK Health Security Agency</td>
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<td>QSAR</td>
<td>Quantitative structure Activity Relationships</td>
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Technical information


High throughput screening (HTS): is a method for scientific experimentation especially used in drug discovery and relevant to the fields of biology and chemistry. Using robotics, data processing/control software, liquid handling devices, and sensitive detectors, high-throughput screening allows a researcher to quickly conduct millions of chemical, genetic, or pharmacological tests. Through this process one can rapidly identify active compounds, antibodies, or genes that modulate a particular biomolecular pathway. The results of these experiments provide starting points for drug design and for understanding the noninteraction or role of a particular location.

Integrated approaches to testing and assessment (IATAs): provide a means by which all relevant and reliable information about a chemical is used to answer a defined hazard characterization question. Information considered can include toxicity data, computational model predictions, exposure routes, use cases, and production volumes. This information is used to characterize outcomes that can inform regulatory decision-making. (Integrated Approaches to Testing and Assessment)

In silico: one performed on computer or via computer simulation.

MANET: is for exploring data, whether raw data, transformed data or model residuals. MANET provides a range of graphical tools specially designed for studying multivariate features. MANET useful for gaining insights into the structure and relationships of their data sets.

Mondrian: is a general-purpose statistical data-visualization system.

Omics: are various disciplines in biology whose names end in the suffix -omics, such as genomics, proteomics, metabolomics, metagenomics and transcriptomics.

Rebuilding a Resilient Britain programme: builds on work to develop government science capability and the external evidence base to support policy development. This report sets out more analysis relating to data and evaluation. It examines existing questions to identify cross-cutting themes, and to provide a platform for engagement between government departments and academics to consider medium and long-term questions.

Spherical cow: is a humorous metaphor for highly simplified scientific models of complex real-life phenomena.
The Fourth Industrial Revolution (4IR): is the fourth major industrial era since the initial Industrial Revolution of the 18th century. It is characterized by a fusion of technologies that is blurring the lines between the physical, digital and biological spheres, collectively referred to as cyber-physical systems.

Zettabytes: are 1,000,000,000,000,000,000 bytes. Zettabyte is approximately equal to a thousand Exabytes, a billion Terabytes, or a trillion Gigabytes.