

2023-08-14

Phenomics as an approach to Comparative Developmental Physiology

McCoy, JCS

<https://pearl.plymouth.ac.uk/handle/10026.1/21748>

10.3389/fphys.2023.1229500

Frontiers in Physiology

Frontiers Media SA

All content in PEARL is protected by copyright law. Author manuscripts are made available in accordance with publisher policies. Please cite only the published version using the details provided on the item record or document. In the absence of an open licence (e.g. Creative Commons), permissions for further reuse of content should be sought from the publisher or author.



OPEN ACCESS

EDITED BY

Robert Roghair,
The University of Iowa, United States

REVIEWED BY

Justin L. Grobe,
Medical College of Wisconsin,
United States
Donna Ann Santillan,
The University of Iowa, United States

*CORRESPONDENCE

Oliver Tills,
✉ oliver.tills@plymouth.ac.uk

RECEIVED 26 May 2023

ACCEPTED 24 July 2023

PUBLISHED 14 August 2023

CITATION

McCoy JCS, Spicer JI, Ibbini Z and Tills O
(2023), Phenomics as an approach to
Comparative Developmental Physiology.
Front. Physiol. 14:1229500.
doi: 10.3389/fphys.2023.1229500

COPYRIGHT

© 2023 McCoy, Spicer, Ibbini and Tills.
This is an open-access article distributed
under the terms of the [Creative
Commons Attribution License \(CC BY\)](#).
The use, distribution or reproduction in
other forums is permitted, provided the
original author(s) and the copyright
owner(s) are credited and that the original
publication in this journal is cited, in
accordance with accepted academic
practice. No use, distribution or
reproduction is permitted which does not
comply with these terms.

Phenomics as an approach to Comparative Developmental Physiology

Jamie C. S. McCoy, John I. Spicer, Ziad Ibbini and Oliver Tills*

School of Biological and Marine Sciences, University of Plymouth, Plymouth, United Kingdom

The dynamic nature of developing organisms and how they function presents both opportunity and challenge to researchers, with significant advances in understanding possible by adopting innovative approaches to their empirical study. The information content of the phenotype during organismal development is arguably greater than at any other life stage, incorporating change at a broad range of temporal, spatial and functional scales and is of broad relevance to a plethora of research questions. Yet, effectively measuring organismal development, and the ontogeny of physiological regulations and functions, and their responses to the environment, remains a significant challenge. “Phenomics”, a global approach to the acquisition of phenotypic data at the scale of the whole organism, is uniquely suited as an approach. In this perspective, we explore the synergies between phenomics and Comparative Developmental Physiology (CDP), a discipline of increasing relevance to understanding sensitivity to drivers of global change. We then identify how organismal development itself provides an excellent model for pushing the boundaries of phenomics, given its inherent complexity, comparably smaller size, relative to adult stages, and the applicability of embryonic development to a broad suite of research questions using a diversity of species. Collection, analysis and interpretation of whole organismal phenotypic data are the largest obstacle to capitalising on phenomics for advancing our understanding of biological systems. We suggest that phenomics within the context of developing organismal form and function could provide an effective scaffold for addressing grand challenges in CDP and phenomics.

KEYWORDS

phenomics, development, bioimaging, embryonic development, Comparative Developmental Physiology

Introduction

It is crucial that we make every effort to identify the grand challenges, the biggest questions in biology. Questions around the development of organismal form and function, and its response to biotic and abiotic factors, presents both opportunity and challenge to researchers, with significant advances in understanding possible through the adoption of innovative approaches to its empirical study. Indeed, our ability to tackle these questions is increasingly reliant on developing technologies for measuring properties of biological systems in new ways (Warburton et al., 2006). The term ‘omics’ is now established in biology as a global approach to measuring biological systems at different levels of organisation, from genomics, transcriptomics and metabolomics at biochemical levels,

through to phenomics - measurement of the phenotype at the level of the whole organism (Houle et al., 2010). ‘Omics’ approaches increase the scale, resolution, or breadth of measurements, often providing a so-called ‘global’ measure of biological response at a particular level of organisation. Omics have become one of the central facilitators of addressing grand challenges in biological research.

The use of omics at the subcellular level is now commonplace in every area of biological research. Adoption of biochemical omics has been fuelled by technological innovation, enabling transferrable and scalable approaches that are broadly accessible. However, despite being recognized as a significant opportunity for advancing biological research in the 21st century (Houle et al., 2010; Kültz et al., 2013), phenomics remains comparatively underutilized outside of research areas focused on human health, such as crop sciences and medicine. The reasons for this hiatus are difficult to pinpoint, but there is a renewed sense that phenomics is an approach whose time has come. Consequently, here we explore the synergies between phenomics and animal development generally, with a particular emphasis on developmental physiology, particularly Comparative Developmental Physiology, or CDP (Warburton and Burggren, 2005; Warburton et al., 2006). We then suggest ways in which taking a phenomics approach to development is beneficial not only for addressing grand challenges in CDP and development generally, but also in the development of phenomics itself as an approach to studying complex biological systems.

What is phenomics?

The phenotype is the ultimate expression of biological organisation at the organismal level. It is also the most visible, striking in its breadth and complexity. Approaches to studying whole-organismal biology typically become operational by restricting what can otherwise be overwhelming complexity, to a relatively small number of observable and tractable traits. Indeed, without such selection, any experimental design rapidly becomes unmanageable. However, the process of deciding what to measure is a key step in experimental design and one that has significant implications for the results and interpretation of any experiment. While we trust that our subject knowledge and biological intuition inform this process, there is undoubtedly a significant element of chance associated with this step of experimental design and one that can affect not only the results of an experiment, but even the trajectory of a researcher’s career (Houle et al., 2010; Lürig et al., 2021).

Here, we define phenomics as “the acquisition of high-dimensional phenotypic data on an organism-wide scale” (*sensu* Houle et al., 2010) and the phenome as “the phenotype of the organism as a whole, including the sum of its morphology, physiology and behaviour” (*sensu* Keller et al., 1992). The term ‘phenome’ was first used by Davis (1949) to describe “the phenotype as a whole”. Along with Werner K. Maas, they coined the term to describe, “the sum of extragenic, non-autoreproductive portions of the cell, whether cytoplasmic or nuclear”. This view was advanced by Soulé (1964) who referred to the phenome as ‘the phenotypic analog of the genome’. Despite these earlier uses of the term, more recent studies frequently assign coinage of the term ‘phenomics’ in

reference to the study of the phenome, to Steven A. Garan in 1996 (Yu and Fang, 2009; Shi et al., 2014; Jin, 2021). Irrespective of its origins, phenomics is currently contextualised as an approach involving the acquisition of phenotypic information at genome wide scales, and it has frequently been regarded as such in reviews and textbooks (Bilder et al., 2009; Furbank 2009; Houle et al., 2010; Hancock, 2014; Tardeiu et al., 2017).

While the notion of phenomics has been around for more than half a century, its study and use as an approach to biological research has only recently gained significant traction. Houle et al. (2010) highlighted its advantages over traditional approaches to studying the phenotype, and others (e.g., Schork, 1997; Freimer and Sabatti, 2003; Bilder et al., 2009) have described it as the natural complement to genomics. Despite this, much like the Human Genome Project, which faced considerable opposition in the face of the counter-claim that reductionist approaches to the study of selected regions of the genome were sufficient for its understanding, phenomics initially met considerable indifference, perhaps suggesting that current approaches to organismal phenotyping were considered sufficient for addressing biological questions of interest (Houle et al., 2010). It is worth noting that while the Human Genome Project fuelled major advancements in biological understanding and acted as the catalyst for advances in modern medicine (Gibbs, 2020), it also met with opposition and some scepticism (Houle et al., 2010; Moraes and Góes, 2016). However, subsequently, molecular omics has necessitated both significant investment and a change in thinking, moving from a reductionist approach to large integrated analyses of complex biological responses, and it has been associated with step-changes in the bioinformatics used for the acquisition and analysis of these data.

Whilst initial traction was lacking in the progression of phenomics as an approach, technological advancements enabling the acquisition of phenotypic data at the whole organism level has accelerated its integration into multiple streams of biological research. Houle et al. (2010) highlighted that limiting the number of phenotypic traits measured to those with some pre-established functional significance could obscure the identification of potentially important traits implicit in the biological response or endpoint of interest, and that high-dimensional phenotyping was necessary to identify the traits, or combinations of traits, that really matter. Now, phenomics is regularly utilised in characterising the genetic basis of complex traits, for tackling disease (Denny et al., 2010; Pendergrass et al., 2011; 2013; Hebring, 2014; Özdemir, 2020), selective breeding (Cossa et al., 2021) and in the characterisation of responses to toxicants (Adira et al., 2020; 2021; Hussain et al., 2020). Phenomics is also becoming increasingly utilised in the assessment of responses to environmental change, particularly within the crop sciences (e.g., Warringer et al., 2003; Schnaubelt et al., 2013; Singh et al., 2018; Adhikari et al., 2019; Li et al., 2019; Marsh et al., 2021; Tills et al., 2021; 2022).

The explosion of literature adopting high-dimensional phenotyping approaches in the crop and medical literature in the last decade was facilitated by the development of transferable technologies for the acquisition of phenotypic data at whole organism scales in these systems (Bilder et al., 2009; Finkel, 2009; Furbank and Tester, 2011; Großkinsky et al., 2015; Tardieu et al., 2017; Grapov et al., 2018; Roitsch et al., 2019; Nabwire et al., 2021). This includes advancements in phenotyping technologies for

common model species including *Arabidopsis* (Furbank and Tester, 2011; Vanhaeren et al., 2015), the zebrafish *Danio rerio* (Xu et al., 2010; Pelkowski et al., 2011; Peravali et al., 2018; Spomer et al., 2012), the nematode *Caenorhabditis elegans* (White et al., 2010; Olmedo et al., 2015) and the fruit fly *Drosophila melanogaster* (Dagani et al., 2007; Chung et al., 2010; Levario et al., 2016). In a practical sense, phenomics typically takes the form of sensors, combined with some degree of automation, whether analytical such as computer vision pipelines (Tills et al., 2018; Choudhury et al., 2019), or physical such as robotics for processing samples (Yang et al., 2020). Consequently, adding dimensionality to the measurement of the phenotype may include greater temporal or spatial resolution, thereby increasing the scale of experiments, or integrating a wider range of sensing modalities and analytical approaches. The solution provided by phenomics is largely driven by the interaction between the biological system and experimental design, and owing to the incredible diversity in what we term the “phenotype”, what phenomics “looks like” can vary considerably. For example, within the crop sciences, phenomics can be executed by robotic systems in greenhouses, and automated tractors and drones out in the field (Tardieu et al., 2017; Zhao et al., 2019; Yang et al., 2020). In both environments, the integration of a wide array of high-throughput sensing modalities is often undertaken, and the resulting data has greater dimensionality than would otherwise be possible.

More recently, phenomics has seen application to address questions relating to animal ecology and evolution, much of which has been facilitated by the development and application of computer vision approaches (reviewed by Lürig et al., 2021). Lürig et al. (2021) provide a review of computer vision approaches and their utility in integrating phenomics into the fields of ecological and environmental research, as well as the general workflows associated with this. Application of computer vision-based approaches alleviate many of the bottlenecks associated with the acquisition and analysis of phenotypic data at large scales. This ability to apply more powerful analyses to answer biological questions is a key strength of phenomics approaches more generally. Advances in this area are also being supported by the use of deep learning (Grapov et al., 2018; Nabwire et al., 2021) and non-linear dimensionality reduction (Tills et al., 2021; Tills et al., 2022), both of which can be powerful enablers for dealing with complex systems. Phenomics approaches are now underpinning applications including the move towards personalised medicine, where large datasets of clinical data for individual patients, from molecular to phenotypic levels of biological organisation, are used to inform treatment decisions (Curcin, 2020).

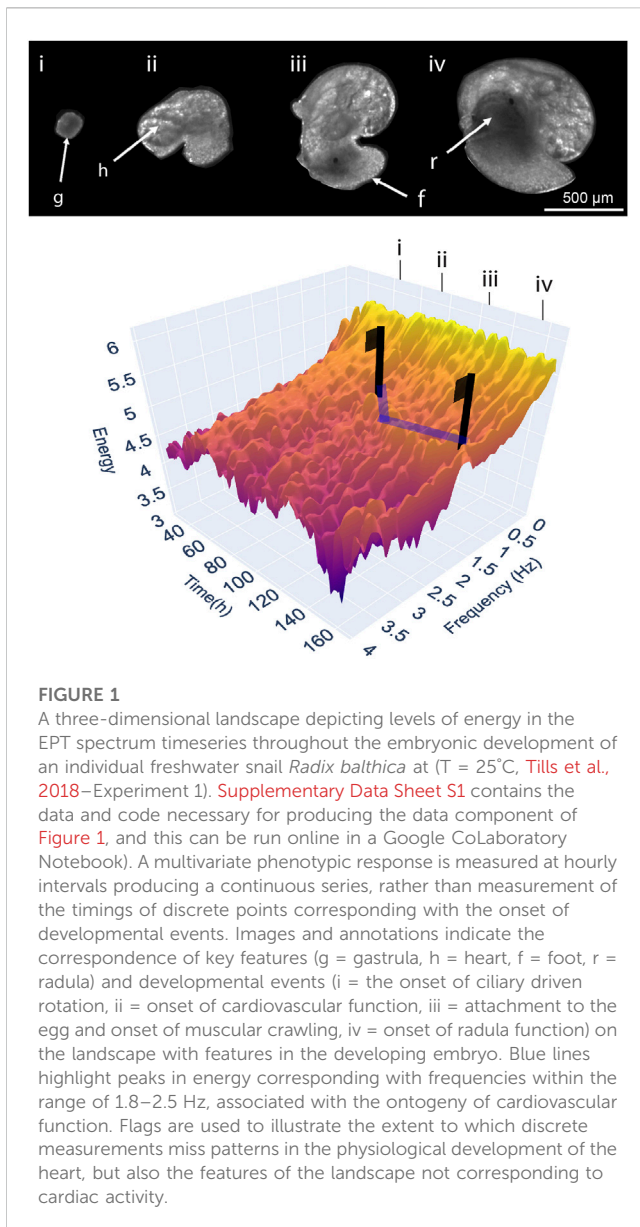
The reasons for phenomics not having been adopted more broadly outside of these areas of research are multifaceted, but likely includes the phenome not being as amenable to transferrable methods as is the case for the subject of molecular omics, combined with a belief that current approaches to phenotyping are adequate. There has been an acknowledged shift (both financial and academic) towards quantifying biological systems at lower levels of biological organisation (Kültz et al., 2013; Noble, 2010), with investment and innovation directed accordingly, towards molecular-omics. Despite the renaissance of whole-organism perspectives on the phenotype in the crop sciences and medicine, it remains the case that integration

of phenomics into research areas outside of these disciplines, particularly to non-model species, remains in its infancy.

Phenomics and Comparative Developmental Physiology

The ability to interrogate an intricate and dynamic biological system is arguably one of the greatest attractions of working with developing organisms, yet it is these same attributes that render their study particularly challenging. Phenomics appears to be a natural complement to this challenge, by enabling the acquisition of phenotypic data at scales that integrate the multifaceted nature of biological responses during this dynamic period of life. Animal development comprises considerable functional and spatial change, with responses to biotic and abiotic factors constituting changes to an array of individual traits. Therefore, using phenomics to study early development brings major advantages, including reducing the element of chance associated with pre-selecting the trait(s) believed to be of functional significance to our biological response of interest. We are not the first to promote a phenomics approach to the study of development—indeed sciomics and high-throughput phenotyping were both proposed in Comparative Developmental Physiology—Contributions, Tools and Trends (Feder, 2006; Spicer, 2006) as being central to the advancement of development, and developmental physiology in particular. Phenomics, however, extends beyond simply widening the dimensionality of phenotypic measurement. Instead, phenomics, and the omics more generally, have the capacity to bring better understanding of biological systems through their analysis in a combinatorial manor, rather than as a sum of their parts. An organism is a functionally integrated unit (Forsman, 2015), and consequently the way in which we measure it should reflect this.

In our own research, we are creating new analytical methods for measuring the phenome of developing organisms. First, as appears to be a frequent precursor to the application of phenomics, we designed and built technologies with which to acquire phenotypic data at whole organism scales, in the form of custom bioimaging hardware—a laboratory instrument for high-throughput and long-term imaging of aquatic embryos (Tills et al., 2018; Tills et al., 2022). The video produced using this approach made immediately clear the limitation of manual approaches to studying a dynamic process. Initially with a focus on developmental events, we were able to observe the complexity surrounding the timing of specific morphological and physiological transitions, such as the onset of cardiovascular function, the initiation of ciliary driven rotation, or more subtle muscular driven behaviours (Tills et al., 2013). Due to the disjunct between the amount of phenotypic information acquired, and what could be feasibly analysed manually by a user, we began considering holistic and non-supervised approaches to measuring development—moving away from trait-specific approaches (Tills et al., 2013; Tills et al., 2022). Most recently, in an attempt to integrate as broad a range of a developing organism’s physiology and behaviour, this has taken the form of an approach termed ‘energy proxy traits’ (Tills et al., 2021). EPTs measure all observable movement-based characteristics of an individual in a video as a spectrum of energy associated with frame-to-frame signals in the brightness of fluctuating mean pixel values. Fluctuations in mean pixel values, the level of brightness in different regions of an image, are the most basic of image statistics. Yet EPTs have proven to be an effective approach to



integrating complex developmental physiology (Tills et al., 2022), but also in quantifying cardiac physiology in a range of different species (Ibbini et al., 2022).

Traditional approaches to measuring development can be likened to erecting flags on a high-dimensional landscape (surface), built from a dataset consisting of development (x -axis), different phenotypic variables (y -axis), and their response (z -axis) each depicted along their own axes (Figure 1). Development, and the ontogeny of physiological traits, are frequently treated as a series of discrete events or stages, despite the process itself being a continuum (Burggren, 2021a; b) and their usage largely ignores any phenotypic transition between these periods (“flags”) in development. This continuous representation of organismal development can be extended, by considering experimental variables such as individual, genotype or experimental treatment as increasingly stacked landscapes, and the differences between them as the net experimental response. Figure 1 portrays such a landscape plotted with real EPT spectra throughout the 7-day embryonic development

of the freshwater gastropod *Radix balthica*. The landscape has been annotated indicating the occurrence of major developmental events and transitions, and with vectors indicating the signal associated with the ontogeny of cardiac physiology. Flags placed on the vectors highlighting the signal associated with cardiac physiology indicate how discrete measurement of heart rate at these timepoints would reflect only a small part of the development of heart rate, but also the developmental physiology of the developing embryo more broadly. Within the EPT landscape are troughs linked to manually identifiable transitions in embryo locomotory behaviours, the ontogeny of whole body and gut muscular contractions, but also an integration of all observable organismal physiology much of which is unquantifiable using individual trait-based quantification. Furthermore, considerable variation is also evident at finer scales between these major behavioural and physiological transitions. This stochasticity highlights the implicit variability of movement associated with embryonic behaviours and physiologies on a timepoint-by-timepoint (hourly) basis. We use the flag-landscape analogy to highlight the major limitation of applying limited discrete measurements to the dynamic and continuous system of development. By measuring the phenotype at specific stages in development, considerable changes in observable phenotype are missed (Burggren, 2021a). We use EPTs as they are an approach to development capable of high-dimensional continuous quantification, but a ‘developmental landscape’ could equally incorporate other physiological measures such as growth curves, metabolic rates, or other physiological rates, or indeed be applied to modelling sensitivity (Burggren and Mueller, 2015). Complex developmental landscapes illustrate the power of Comparative Developmental Physiology to contribute to the growth of phenomics as an approach, and to the phenome as a concept.

What can we learn from where phenomics is established?

Phenomics is most established in research areas where there is a short-term human-benefit to the systematic and high-throughput study of the phenotype, including the selective breeding of crops (Furbank, 2009; Großkinsky et al., 2015; Neto and Borém, 2015; Lobos et al., 2017; Borrill et al., 2018; York, 2019; Yang et al., 2020; Razzaq et al., 2021) and livestock (Greenwood et al., 2015; Cole et al., 2020; Pérez-Enciso and Stiebel, 2021), biosciences (Čapek et al., 2023) and medicine (Denny et al., 2010; Pendergrass et al., 2011; 2013; Maslov et al., 2018; Özdemir, 2020). Each of these research areas has achieved a technological capacity to study the phenotype with a level of throughput and thoroughness that enables the acquisition of phenome level data; however, subsequent analysis of data at these scales also presents significant challenges. Within the context of animal development, this challenge is magnified given the often considerable functional and morphological changes associated with embryonic and larval development. What then, can we learn from other areas of research that integrate similarly high-dimensional datasets?

Phenomics produces high-dimensional datasets with potentially thousands of predictor-outcome variable relationships (Furbank, 2009; Houle et al., 2010), and its integration into animal development and comparative developmental physiology is thereby limited as much by our ability to analyse and handle data at such scales, as the collection of data themselves (Tardieu et al., 2017). Whilst this is not the place to review the challenges associated with

analysing data at these scales, and the statistical techniques used to navigate these challenges, we direct readers to [Houle et al. \(2010\)](#) and [Hebbring \(2014\)](#) for detailed discussion of each of these points. Within the molecular omics, there is fierce and entrenched debate amongst biologists about the potential pitfalls of high-dimensional datasets, and so it seems appropriate to consider these here. Omics datasets can be likened to the mythical Pandora's box, in that whilst they enable measurement of biology at an unprecedented scale, they also produce vast datasets that can require considerable time and expertise to interpret and can support research pre-hypotheses-testing. Advancement of bioinformatics approaches has undoubtedly been central to the analysis and interpretation of these data, enabling analyses of large omics datasets for both data exploration and hypothesis testing. Integration of phenomics into areas of research including animal development and developmental physiology will therefore necessitate drawing on expertise in bioinformatics and computational biology, all geared towards addressing challenges associated with the acquisition, analysis and storage of data, including handling metadata, ensuring data reproducibility, and driving data standards. As phenomics grows as an approach, and as it is increasingly integrated into a wider diversity of research areas, it becomes increasingly important to learn these lessons from molecular bioinformatics.

What can Comparative Developmental Physiology contribute to phenomics?

Phenomics as an approach will undoubtedly reward researchers working in developmental biology in their ability to address challenging research questions that capitalise on the breadth and integrated nature of this period of life. However, we also suggest morphological differentiation and physiological ontogeny inherent in organismal development as a model system, has significant potential to develop phenomics as an approach, contributing to advancing many areas of biological research including those identified in [Burggren's \(2021a\)](#) "Developmental Physiology: Grand Challenges".

Organismal development is inherently complex, multifaceted and dynamic. Owing to the small size of early life stages and their taxonomic universality, animals during their early development present an excellent model for the application of phenomics, both in a practical sense, and in their capacity to stretch and develop the theoretical understanding surrounding phenomics. Embryos are, after all, used in research ranging from environmental science and ecotoxicology, through to drug discovery and evolutionary developmental biology. The phenome is highly heterogeneous in the types of data required to describe it, and therefore the methods and approaches necessary to measure it. This contrasts markedly with the universality of molecular-omics, with transferrable approaches for both data acquisition and analysis, but also an accessible shared understanding of what each of these molecular-omics are across all areas of biology. Our knowledge of the phenome is crude, and poorly endowed with transferrable approaches compared with the other omics. We therefore support and emphasise the points made by others, that phenomics can act as a key enabler to the advancement of science in the 21st century.

While the potential contribution of embryonic development to advancing the concept of the phenome is multifaceted, we identify four key strengths of developing embryos as models for phenomics:

- i) High-dimensional phenotyping of embryos is highly scalable, owing to their small size and ability to be cultured within a laboratory setting. This scalability can enable high-throughput phenotyping approaches of complex phenotypes, with associated large-scale data collection.
- ii) Biodiversity, including physiological diversity, mean that application of phenomics to embryonic development not only presents opportunities to advance our understanding of the phenome of this critical period of life across the natural world, but to also accelerate the development of approaches used to measure and analyse it, by drawing on this biodiversity.
- iii) Central to phenomics is high-dimensional phenotyping. The information content of the phenome during embryonic development is extremely high (and perhaps greater than at any other life stage), given the considerable temporal, spatial and functional change associated with this period of life. Static trait-based approaches are therefore a well-recognised limitation to its study. Consequently, advancement of technological, theoretical, and analytical approaches to enable the application of phenomics will provide significant opportunity to further our understanding of these dynamic biological systems.
- iv) Research to understand the role of embryonic development in evolutionary and ecological processes is broad, and so too are the associated frameworks, models and approaches created to support it. These offer considerable opportunities for the advancement of phenomics, by acting as a scaffold from which to build and interpret new approaches to understanding the phenotype.

Priorities and future directions

The current pace of technological innovation and change is unparalleled. Consequently the "art of the possible" is rapidly expanding. Undoubtedly this presents significant opportunities to the advancement of phenomics as an approach. However, to capitalise on technological advances it requires a parallel initiative advancing the development of methods, adoption of technologies and training of staff that span both areas. These activities have been central to the advancement of molecular-omics. The pace of technological innovation and the general lack of accessible commercial products in the life sciences has positioned open-source DIY approaches to become common enablers of innovation and research using phenomics. Key strengths of open-source include greater accessibility, accelerated innovation and increased return on investment, but open-source is certainly not a panacea with prerequisite skills, equipment and resources, acting as a potential barrier to adoption. Technological innovation is fuelling the acquisition of phenotypic data at unprecedented scales ([Meijering et al., 2016](#)), with bioimaging being one of the greatest contributors to this growth. Here, we have made the case for embryonic development as a model for phenomics, with a focus on how the benefits will be reciprocal. The application of phenomics as an approach, we believe, can lead to significant advancements in our understanding of the evolutionary and ecological significance of changes in the phenotype during early development. In doing so, phenomics as an approach will itself be advanced by using this complex, dynamic and scalable biological system.

The challenge of phenomics is particularly timely given our global biodiversity crisis adding considerable urgency to assessment of biological sensitivity and other conservation physiology themed research activities. Current capacity for phenomics is largely limited to frequently used model species, likely due to the considerable resources associated with developing computational tools capable of measuring high-dimensional phenotypic change (Lussier and Liu, 2007; Xu et al., 2010; Perivali et al., 2018; Olmedo et al., 2015; Lürig et al., 2021). As it stands, phenomics is largely not transferrable taxonomically, regardless of the life stage. As a result, the capacity for phenomics to contribute to research geared towards conservation of biodiversity is currently limited. Comparative Physiology and Comparative Development Physiology operates on the basis of the Krogh principle (Krogh, 1929), in which species are selected based on their suitability to answer the biological question of interest. Consequently, development of high-dimensional phenotyping approaches with levels of transferability comparable to those of the molecular omics should be a priority if we are to facilitate the integration of phenomics into the study of animal development, outside of model species.

In summary, phenomics as an approach provides exciting opportunities for advancing our understanding of the evolutionary and ecological significance of developmental change, the genetic underpinnings of alterations to early development, as well as in interpreting developmental responses to biotic and abiotic change. Directing future research towards the establishment of high-dimensional phenotyping approaches that have the broadest relevance and applicability across the animal kingdom should be a key priority, and here, we identify animal development, particularly developmental physiology, as having significant potential to catalyse this endeavour.

Data availability statement

The original contributions presented in the study are included in the article/[Supplementary Material](#), further inquiries can be directed to the corresponding author.

References

- Adhikari, N. D., Simko, I., and Mou, B. (2019). Phenomic and physiological analysis of salinity effects on lettuce. *Sensors* 19, 4814. doi:10.3390/s19214814
- Audira, G., Lai, Y. H., Huang, J. C., Chen, K. H. C., and Hsiao, C. D. (2021). Phenomics approach to investigate behavioral toxicity of environmental or occupational toxicants in adult zebrafish (*Danio rerio*). *Curr. Protoc.* 1, e223. doi:10.1002/cpz1.223
- Audira, G., Siregar, P., Chen, J. R., Lai, Y. H., Huang, J. C., and Hsiao, C. D. (2020). Systematical exploration of the common solvent toxicity at whole organism level by behavioral phenomics in adult zebrafish. *Environ. Pollut.* 266, 115239. doi:10.1016/j.envpol.2020.115239
- Bilder, R. M., Sabb, F. W., Cannon, T. D., London, E. D., Jentsch, J. D., Parker, D. S., et al. (2009). Phenomics: the systematic study of phenotypes on a genome wide scale. *Neuroscience* 164, 30–42. doi:10.1016/j.neuroscience.2009.01.027
- Borrill, P., Harrington, S. A., and Uauy, C. (2018). Applying the latest advances in genomics and phenomics for trait discovery in polyploid wheat. *Plant J.* 97, 56–72. doi:10.1111/tpj.14150
- Burggren, W. W. (2021a). Developmental physiology: grand challenges. *Front. Physiology* 12, 706061. doi:10.3389/fphys.2021.706061
- Burggren, W. W., and Mueller, C. A. (2015). Developmental critical windows and sensitive periods as three-dimensional constructs in time and space. *Physiological Biochem. Zoology* 88, 91–102. doi:10.1086/679906
- Burggren, W. W. (2021b). Putting the August Krogh principle to work in developmental physiology. *Comp. Biochem. Physiology -Part A Mol. Integr. Physiology* 252, 110825. doi:10.1016/j.cbpa.2020.110825
- Čapek, D., Safroshkin, M., Morales-Navarrate, H., Toulany, N., Arutyunov, G., Kurzbach, A., et al. (2023). EmbryoNet: using deep learning to link embryonic phenotypes to signaling pathways. *Nat. Methods*, 1–9.
- Choudhury, S. D., Samal, A., and Awada, T. (2019). Leveraging image analysis for high-throughput plant phenotyping. *Front. Plant Sci.* 10, 2080–2088. doi:10.3389/fpls.2019.00508
- Chung, K., Kim, Y., Gong, E., Shvartsman, S., and Lu, H. (2010). “High-throughput end-on imaging of drosophila embryo for quantitative analysis of morphogens and signalling,” in 14th International conference on Miniaturized Systems for Chemistry and Life Sciences, Groningen, The Netherlands, 3–7 October–2010.
- Cole, J. B., Eaglan, S. A. E., Maltecca, C., Mulder, H. A., and Pryce, J. E. (2020). The future of phenomics in dairy cattle breeding. *Anim. Front.* 10, 37–44. doi:10.1093/af/vfaa007
- Crossa, J., Fritsche-Neto, R., Montesinos-Lopez, O. A., Costa-Neto, G., Dreisigacker, S., Montesinos-Lopez, A., et al. (2021). The modern plant breeding triangle: optimizing the use of genomics, phenomics, and enviromics data. *Front. Plant Sci.* 12, 651480. doi:10.3389/fpls.2021.651480
- Curcin, Y. (2020). Why does human phenomics matter today? *Learn. Health Syst.* 4, e10249. doi:10.1002/lrh2.10249
- Dagani, G. T., Monzo, K., Fakhoury, J. R., Chen, C. C., Sisson, J. C., and Zhang, X. (2007). Microfluidic self-assembly of live *Drosophila* embryos for versatile high-throughput analysis of embryonic morphogenesis. *Biomed. Microdevices* 9, 681–694. doi:10.1007/s10544-007-9077-z

Author contributions

All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication.

Funding

This project was supported by a UKRI Future Leaders Fellowship to OT (MR/T01962X/1), BBSRC Follow on Fund to OT, JS and JM (BB/W017938/1), a NERC ARIES DTP studentship to JM and a University of Plymouth PhD studentship to ZI.

Conflict of interest

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

Publisher's note

All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Supplementary material

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fphys.2023.1229500/full#supplementary-material>

- Davis, B. D. (1949). The isolation of biochemically deficient mutants of Bacteria by means of penicillin. *Proc. Natl. Acad. Sci. U. S. A.* 35, 1–10. doi:10.1073/pnas.35.1.1
- Denny, J. C., Ritchie, M. D., Basford, M. A., Pulley, J. M., Bastarache, L., Brown-Gentry, K., et al. (2010). PheWAS: demonstrating the feasibility of a phenome-wide scan to discover gene-disease associations. *Bioinformatics* 26, 1205–1210. doi:10.1093/bioinformatics/btq126
- Feder, M. E. (2006). “Sciomics: community/model organism based and individualistic research strategies for comparative animal developmental physiology,” in *Comparative developmental physiology: Contributions, tools and trends*. Editors S. J. Warburton, W. W. Burggren, B. P. Pelster, C. L. Reiber, and J. I. Spicer (New York: Oxford University Press), 161–173.
- Finkel, E. (2009). Imaging. With ‘phenomics,’ plant scientists hope to shift breeding into overdrive. *Science* 325, 380–381. doi:10.1126/science.325_380
- Forsman, A. (2015). Rethinking phenotypic plasticity and its consequences for individuals, populations and species. *Heredity* 115, 276–284. doi:10.1038/hdy.2014.92
- Freimer, N., and Sabatti, C. (2003). The human phenome Project. *Nat. Genet.* 34, 15–21. doi:10.1038/ng0503-15
- Furbank, R. T. (2009). Plant phenomics: from gene to form and function. *Funct. Biol.* 36, 5–6. doi:10.1016/j.fplants.2011.09.005
- Furbank, R. T., and Tester, M. (2011). Phenomics - technologies to relieve the phenotyping bottleneck. *Trends Plant Sci.* 16, 635–644. doi:10.1016/j.fplants.2011.09.005
- Gibbs, R. A. (2020). The human genome Project changed everything. *Nat. Rev. Genet.* 21, 575–576. doi:10.1038/s41576-020-0275-3
- Grapov, D., Fahrman, J., Wanichthanarak, K., and Khoomrung, S. (2018). Rise of deep learning for genomic, proteomic, and metabolomic data integration in precision medicine. *Omics A J. Integr. Biol.* 22, 630–636. doi:10.1089/omi.2018.0097
- Greenwood, P. L., Bishop-Hurley, G. J., González, L. A., and Ingham, A. B. (2015). Development and application of a livestock phenomics platform to enhance productivity and efficiency at pasture. *Animal Prod. Sci.* 56, 1299–1311. doi:10.1071/an15400
- Großkinsky, D. K., Svendsgaard, J., Christensen, S., and Roitsch, T. (2015). Plant phenomics and the need for physiological phenotyping across scales to narrow the genotype-to-phenotype knowledge gap. *J. Exp. Bot.* 66, 5429–5440. doi:10.1093/jxb/erv345
- Hancock, J. M. (2014). *Phenomics*. New York: CRC Press.
- Hebbring, S. J. (2014). The challenges, advantages and future of phenome-wide association studies. *Immunology* 141, 157–165. doi:10.1111/imm.12195
- Houle, D., Govindaraju, D. R., and Omholt, S. (2010). Phenomics: the next challenge. *Nat. Rev. Genet.* 11, 855–866. doi:10.1038/nrg2897
- Hussain, A., Audira, G., Malhotra, N., Uapipatanakul, B., Chen, J. R., Lai, Y. H., et al. (2020). Multiple screening of pesticides toxicity in zebrafish and daphnia based on locomotor activity alterations. *Biomolecules* 10, 1224. doi:10.3390/biom10091224
- Ibbini, Z., Spicer, J. I., Truebano, M., Bishop, J., and Tills, O. (2022). HeartCV: a tool for transferrable, automated measurement of heart rate and heart rate variability in transparent animals. *J. Exp. Biol.* 225, jeb244729. doi:10.1242/jeb.244729
- Jin, L. (2021). Welcome to the phenomics journal. *Phenomics* 1, 1–2. doi:10.1007/s43657-020-00009-4
- Keller, E., Lloyd, E., and Lewontin, R. C. (1992). “Genotype and phenotype,” in *Keywords in evolutionary biology*. Editors E. Keller and E. Lloyd (Cambridge, MA: Harvard University Press), 26–33.
- Krogh, A. (1929). The progress of physiology. *Am. J. Physiology* 90, 243–251. doi:10.1152/ajplegacy.1929.90.2.243
- Kültz, D., Clayton, D. F., Robinson, G. E., Albertson, C., Carey, H. V., Cummings, M. E., et al. (2013). New frontiers for organismal biology. *BioScience* 63, 464–471. doi:10.1525/bio.2013.63.6.8
- Levario, T. J., Zhao, C., Rouse, T., Shvartsman, S. Y., and Lu, H. (2016). An integrated platform for large-scale data collection and precise perturbation of live *Drosophila* embryos. *Sci. Rep.* 6, 21366. doi:10.1038/srep21366
- Li, X., Kong, H., Ji, X., Gao, Y., and Jin, M. (2019). Zebrafish behavioral phenomics applied for phenotyping aquatic neurotoxicity induced by lead contaminants of environmentally relevant level. *Chemosphere* 224, 445–454. doi:10.1016/j.chemosphere.2019.02.174
- Lobos, G. A., Camargo, A. V., del Pozo, A., Araus, J. L., Ortiz, R., and Doonan, J. H. (2017). Editorial: plant phenotyping and phenomics for plant breeding. *Plant Sci.* 8, 2181. doi:10.3389/fpls.2017.02181
- Lürig, M. D., Donoughe, S., Svensson, E. I., Porto, A., and Tsuboi, M. (2021). Computer vision, Machine learning, and the promise of phenomics in ecology and evolutionary biology. *Front. Ecol. Evol.* 09, 642774. doi:10.3389/fevo.2021.642774
- Lussier, Y. A., and Liu, Y. (2007). Computational approaches to phenotyping: high-throughput phenomics. *Proc. Am. Thorac. Soc.* 4, 18–25. doi:10.1513/pats.200607-142JG
- Marsh, J. I., Hu, H., Gill, M., Batley, J., and Edwards, D. (2021). Crop breeding for a changing climate: integrating phenomics and genomics with bioinformatics. *Theor. Appl. Genet.* 134, 1677–1690. doi:10.1007/s00122-021-03820-3
- Maslov, D. L., Balashova, E. E., Trifonova, O. P., and Likhov, P. G. (2018). Metabolomics-based approach to pharmacotherapy personalization: advantages and limitations. *Advantages limitations Curr. Pharmacogenomics Personalized Med.* 16, 192–198. doi:10.2174/1875692116666181008144905
- Meijering, E., Carpenter, A. E., Peng, H., Hamprecht, F. A., and Olivo-Marin, J. C. (2016). Imagining the future of bioimage analysis. *Nat. Biotechnol.* 34, 1250–1255. doi:10.1038/nbt.3722
- Moraes, F., and Góes, A. (2016). A decade of human genome project conclusion: scientific diffusion about our genome knowledge. *Biochem. Mol. Biol. Educ.* 44, 215–223. doi:10.1002/bmb.20952
- Nabwire, S., Suh, H. K., Kim, M. S., Baek, I., and Cho, B. K. (2021). Review: application of artificial intelligence in phenomics. *Sensors* 21, 4363. doi:10.3390/s21134363
- Neto, F., and Borém, A. (2015). *Phenomics: How next-generation phenotyping is revolutionizing plant breeding*. New York: Springer Publishing.
- Noble, D. (2010). Biophysics and systems biology. *Philosophical Trans. Ser. A, Math. Phys. Eng. Sci.* 368, 1125–1139. doi:10.1098/rsta.2009.0245
- Olmedo, M., Geibel, M., Artal-Sanz, M., and Merrow, M. (2015). A high-throughput method for the analysis of larval developmental phenotypes in *Caenorhabditis elegans*. *Genetics* 201, 443–448. doi:10.1534/genetics.115.179242
- Özdemir, V. (2020). Phenomics 2.0: real-world real-time patient outcomes measured by the internet of pharmaceutical things. *OMICS A J. Integr. Biol.* 24, 119–121. doi:10.1089/omi.2020.0015
- Pelkowski, S. D., Kapoor, M., Richendrer, H. A., Wang, X., Colwill, R. M., and Creton, R. (2011). A novel high-throughput imaging system for automated analyses of avoidance behavior in zebrafish larvae. *Behav. Brain Res.* 223, 135–144. doi:10.1016/j.bbr.2011.04.033
- Pendergrass, S. A., Brown-Gentry, K., Dudek, S., Frase, A., Torstenson, E. S., Goodloe, R., et al. (2013). Phenome-wide association study (PheWAS) for detection of pleiotropy within the population architecture using genomics and epidemiology (PAGE) network. *PLoS Genet.* 9, e1003087. doi:10.1371/journal.pgen.1003087
- Pendergrass, S. A., Brown-Gentry, K., Dudek, S. M., Torstenson, E. S., Ambite, J. L., Avery, C. L., et al. (2011). The use of phenome-wide association studies (PheWAS) for exploration of novel genotype-phenotype relationships and pleiotropy discovery. *Genet. Epidemiol.* 35, 410–422. doi:10.1002/gepi.20589
- Peravali, R., Gehrig, J., Giselbrecht, S., Lütjohann, D. S., Hadzhev, Y., Müller, F., et al. (2018). Automated feature detection and imaging for high-resolution screening of zebrafish embryos. *BioTechniques* 50, 319–324. doi:10.2144/000113669
- Pérez-Enciso, M., and Steibel, J. P. (2021). Phenomes: the current frontier in animal breeding. *Genet. Sel. Evol.* 53, 22. doi:10.1186/s12711-021-00618-1
- Razaq, A., Kaur, P., Akhter, N., Wani, S. H., and Saleem, F. (2021). Next-generation breeding strategies for climate-ready crops. *Front. Plant Sci.* 12, 620420. doi:10.3389/fpls.2021.620420
- Roitsch, T., Cabrera-Bosquet, L., Fournier, A., Ghamkar, K., Jiménez-Berni, J., Pinto, F., et al. (2019). Review: new sensors and data-driven approaches—a path to next generation phenomics. *Plant Sci.* 282, 2–10. doi:10.1016/j.plantsci.2019.01.011
- Schnaubelt, D., Schulz, P., Hannah, M. A., Yocgo, R. E., and Foyer, C. H. (2013). A phenomics approach to the analysis of the influence of glutathione on leaf area and abiotic stress tolerance in *Arabidopsis thaliana*. *Front. Plant Sci.* 4, 416. doi:10.3389/fpls.2013.00416
- Schork, N. J. (1997). Genetics of complex disease: approaches, problems, and solutions. *Am. J. Respir. Crit. Care Med.* 156, 103–109. doi:10.1164/ajrccm.156.4.12-tac-5
- Shi, Q., Gao, K., Zhao, H., Wang, J., Zhai, X., Lu, P., et al. (2014). Phenomics research on coronary heart disease based on human phenotype ontology. *BioMed Res. Int.* 2014, 240284. doi:10.1155/2014/240284
- Singh, B., Mishra, S., Bohra, A., Joshi, R., and Siddique, K. H. M. (2018). “Crop phenomics for abiotic stress tolerance in crop plants,” in *Biochemical, physiological and molecular avenues for combating abiotic stress in plants*. Editor S. H. Wani (London: Elsevier), 277–296.
- Soulé, M. E. (1964). *The evolution and population genetics of the side-blotched lizards (Uta stansburiana and relatives) on the islands in the Gulf of California*. Mexico: Stanford University.
- Spicer, J. I. (2006). “A physiological approach to heterochrony,” in *Comparative developmental physiology: Contributions, tools and trends*. Editors S. J. Warburton, W. W. Burggren, B. Pelster, C. L. Reiber, and J. I. Spicer (Oxford: Oxford University Press), 191–202.
- Spomer, W., Pfriem, A., Alshut, R., Just, S., and Pylatiuk, C. (2012). High-throughput screening of zebrafish embryos using automated heart detection and imaging. *J. Laboratory Automation* 17, 435–442. doi:10.1177/2211068212464223

- Tardieu, F., Cabrera-Bosquet, L., Pridmore, T., and Bennett, M. (2017). Plant phenomics, from sensors to knowledge. *Curr. Biol.* 27, R770-R783–R783. doi:10.1016/j.cub.2017.05.055
- Tills, O., Holmes, L. A., Quinn, E., Everett, T., Truebano, M., and Spicer, J. I. (2022). Phenomics enables measurement of complex responses of developing animals to global environmental drivers. *Sci. Total Environ.* 858, 159555. doi:10.1016/j.scitotenv.2022.159555
- Tills, O., Rundle, S. D., and Spicer, J. I. (2013). Variance in developmental event timing is greatest at low biological levels: implications for heterochrony. *Biol. J. Linn. Soc.* 110, 581–590. doi:10.1111/bij.12158
- Tills, O., Spicer, J. I., Grimmer, A., Marini, S., Jie, V. W., Tully, E., et al. (2018). A high-throughput and open-source platform for embryo phenomics. *PLoS Biol.* 16, e3000074. doi:10.1371/journal.pbio.3000074
- Tills, O., Spicer, J. I., Ibbini, Z., and Rundle, S. D. (2021). Spectral phenotyping of embryonic development reveals integrative thermodynamic responses. *BMC Bioinforma.* 22, 232. doi:10.1186/s12859-021-04152-1
- Vanhaeren, H., Gonzalez, N., and Inzé, D. (2015). A journey through a leaf: phenomics analysis of leaf growth in *Arabidopsis thaliana*. *Arabidopsis Book* 13, e0181. doi:10.1199/tab.0181
- Warburton, S. J., and Burggren, W. W. (2005). Comparative developmental physiology: an interdisciplinary convergence. *Annu. Rev. Physiology* 67, 203–223. doi:10.1146/annurev.physiol.67.040403.104223
- Warburton, S. J., Burggren, W. W., Pelster, B., Reiber, C. L., and Spicer, J. I. (2006). *Comparative developmental physiology*. New York: Oxford University Press.
- Warringer, J., Ericson, E., Fernandez, L., Nerman, O., and Blomberg, A. (2003). High-resolution yeast phenomics resolves different physiological features in the saline response. *Proc. Natl. Acad. Sci. U. S. A.* 100, 15724–15729. doi:10.1073/pnas.2435976100
- White, A. G., Cipriani, P. G., Kao, H. L., Lees, B., Geiger, D., Sontag, E., et al. (2010). “Rapid and accurate developmental stage recognition of *C. elegans* from high throughput image data,” in Conference on Computer Vision and Pattern Recognition (CVPR), San Francisco, 13–18 June–2010.
- Xu, X., Xu, X., Huang, X., Xia, W., and Xia, S. (2010). A high-throughput analysis method to detect regions of interest and quantify zebrafish embryo images. *J. Biomol. Screen.* 15, 1152–1159. doi:10.1177/1087057110379155
- Yang, W., Feng, H., Zhang, X., Zhang, J., Doonan, J. H., Batchelor, W. D., et al. (2020). Crop phenomics and high-throughput phenotyping: past decades, current challenges, and future perspectives. *Mol. Plant* 13, 187–214. doi:10.1016/j.molp.2020.01.008
- York, L. M. (2019). Functional phenomics: an emerging field integrating high-throughput phenotyping, physiology, and bioinformatics. *J. Exp. Biol.* 70, 379–386. doi:10.1093/jxb/ery379
- Yu, G., and Fang, X. (2009). Concept of phenomics and its development in plant science. *Mol. Plant Breed.* 7, 639–645.
- Zhao, C., Zhang, Y., Du, J., Guo, X., Wen, W., Gu, S., et al. (2019). Crop phenomics: current status and perspectives. *Front. Plant Sci.* 10, 714. doi:10.3389/fpls.2019.00714