Review Article

**Omics-based ecosurveillance for the assessment of ecosystem function, health, and resilience**

David J. Beale, Oliver A.H. Jones, Utpal Bose, James A. Broadbent, Thomas K. Walsh, Jodie van de Kamp and Andrew Bissett

Land and Water, Commonwealth Scientific and Industrial Research Organisation, Ecosciences Precinct, Dutton Park QLD 4102, Australia; Australian Centre for Research on Separation Science (ACROSS), School of Science, RMIT University, Bundoora West Campus, PO Box 71, Bundoora, VIC 3083, Australia; Agriculture and Food, Commonwealth Scientific and Industrial Research Organisation, Queensland Bioscience Precinct, St Lucia, QLD 4067, Australia; Land and Water, Commonwealth Scientific and Industrial Research Organisation, Acton, ACT 2601, Australia; Oceans and Atmosphere, Commonwealth Scientific and Industrial Research Organisation, Battery Point, TAS 7004, Australia

Correspondence: David J. Beale (david.beale@csiro.au)

Current environmental monitoring efforts often focus on known, regulated contaminants ignoring the potential effects of unmeasured compounds and/or environmental factors. These specific, targeted approaches lack broader environmental information and understanding, hindering effective environmental management and policy. Switching to comprehensive, untargeted monitoring of contaminants, organism health, and environmental factors, such as nutrients, temperature, and pH, would provide more effective monitoring with a likely concomitant increase in environmental health. However, even this method would not capture subtle biochemical changes in organisms induced by chronic toxicant exposure. Ecosurveillance is the systematic collection, analysis, and interpretation of ecosystem health-related data that can address this knowledge gap and provide much-needed additional lines of evidence to environmental monitoring programs. Its use would therefore be of great benefit to environmental management and assessment. Unfortunately, the science of ‘ecosurveillance’, especially omics-based ecosurveillance is not well known. Here, we give an overview of this emerging area and show how it has been beneficially applied in a range of systems. We anticipate this review to be a starting point for further efforts to improve environmental monitoring via the integration of comprehensive chemical assessments and molecular biology-based approaches. Bringing multiple levels of omics technology-based assessment together into a systems-wide surveillance approach will bring a greater understanding of the environment, particularly the microbial communities upon which we ultimately rely to remediate perturbed ecosystems.

**Introduction**

Ecosurveillance, in the context of this review, is defined as the systematic collection, analysis, and interpretation of information on ecosystem health. It synthesises numerous data sources to inform environmental management and policy decisions. Environmental monitoring refers to the measurement of specific parameters, such as nutrients, dissolved oxygen, specific contaminants, and indicator organisms, usually concerning regulatory targets. Our ability to collect and interpret both ecosystem data and the inorganic/organic contaminant information represents a powerful tool for managing environmental health. Ecosurveillance can then also be used to measure the success or failure of interventions in a quantitative manner. This is graphically summarised in Figure 1.

The key point of differentiation between health/surveillance-based measurements of the ecosystem and monitoring of discrete and individual parameters/specific contaminants is that monitoring is more concerned with source-dictated (targeted) measurements (i.e. contaminants, nutrients, etc.) that are present in an environment, whereas ecosystem health metrics are focused on system-wide performance and characterisation (i.e. ecosystem service function/dysfunction due to biotic or abiotic...
perturbation). Omics-based technologies (i.e. genomics, transcriptomics, proteomics, lipidomics, and metabolomics) may be deployed within the ecosurveillance monitoring framework to capture the biological response of the system under perturbation.

Environmental monitoring

A common approach for environmental monitoring is to take a targeted risk-based approach that considers known contaminant sources and associated industrial activities (both historical and current) in the catchment to compile a list of possible contaminants and physical parameters for monitoring [1]. The risk of this approach is that it can only monitor contaminants known to be present, it cannot identify unknown compounds or contaminants of concern that were not known to present, nor consider the cumulative effects of multiple sublethal stressors [2]. It is a static measurement of a fixed set of contaminants at a particular point in time, with no real link to the surrounding system outside of the matrix that was measured. Such assessments, therefore, do not indicate the bioavailability, stability, turnover, or distribution of the substances within the system. This can be a particular problem for contaminants such as perfluoroalkyl substances (PFAS), which are numerous (many undocumented), highly mobile, and are known to bioaccumulate in multiple trophic levels [3,4]; for long-lived contaminants that can bind to and release from sediments (e.g. metals); or, for contaminants that are formed in the system itself (e.g. photochemically produced reactive oxygen species) [5]. Furthermore, understanding the complex interactions between biological systems and anthropogenic environmental changes and the effects of natural variability in weather, season, and diurnal and biogeochemical cycles remains a major research challenge [6–8].

Traditional environmental and organism health monitoring techniques (e.g. chemical monitoring and bioassays) are highly suited to assessing acute toxic effects. They struggle, however, to detect subtle shifts in ecosystem function, species abundances, and animal physiology resulting from low-level and chronic exposures due to changing environmental/climatic conditions and interactions thereof.

Policy decisions are thus often based not on the actual health of the ecosystem, but the quantification of specific contaminants or nutrients above or below an arbitrary legislative level. To some degree, the importance of this gap between the measurement of a substance and the impact will vary with specific circumstances (e.g.
We suggest the data suitable for determining in situ risks of adverse effects or ecosystem perturbation are currently lacking for most environmental contaminants, both for prospective and retrospective assessments. Compounding this further is the uncertainty in ecological risk assessments of chemical mixtures (PFAS, pharmaceuticals, pesticides, chlorinated paraffins, and antibiotics are all cases in point) [10,11]. There is a clear need for better predictive tools for chemical mixture assessment, and the development of in vitro, and in vivo methods to efficiently assess and predict biological effects [12–14]. Effects Directed Analysis (EDA) is one approach to meeting this need, using metabolic sample fractionation, bioassay endpoints, and chemical contaminant analyses to identify key toxicants in the environment [15–17]. While EDA approaches are not the focus of this review, it is acknowledged that they would complement functional omics assays (i.e. metabolomics, lipidomics, and proteomics) used within the ecosurveillance framework proposed within that are aimed at capturing biological information from a sample matrix, which could be further analysed via the EDA pipeline of assays. Furthermore, novel non-targeted screening approaches that can capture contaminants outside of a risk-based monitoring framework have the potential to improve biological contaminant impact assessments and facilitate retrospective data mining for further insights into key contaminants of concern [18], if analysed with appropriately matched omics data (e.g. genomics, transcriptomics, proteomics, or metabolomics).

Ecosurveillance has the potential to bridge the gap in current monitoring data, and link system performance metrics that are tied to measured biotic and abiotic perturbations [19–21]. Omics-based approaches, either applied individually or in combination, have successfully been applied to investigate (potential or measured) function across the tree of life from microbial communities (i.e. microbiomes) through to larger metazoan taxa. Coupling measurable metabolic endpoints to multiple (sometimes subacute) contaminant levels and sources that are currently not identified using a conventional risk-based monitoring program has the potential to provide a more detailed and holistic understanding of temporal, spatial, and multigenerational impacts of contaminants on a single organism or species through to a whole-of-system approach [2]. For example, if one could measure metabolite levels and toxicant concentrations in the same analytical run there would be great potential for directly correlating metabolic changes and the contaminant concentrations needed to cause such changes. Even if contaminant levels were low, the presence of a contaminant, or more likely contaminants, known to cause a particular physiological change before significant harm occurring might at least be used as a trigger for further intensified monitoring and/or remediation before serious harm occurred.

The purpose of this review is to provide an overview of some of the salient applications of environmental functional omics within an omics-based ecosurveillance framework. We also demonstrate how these methods can be coupled with current monitoring data to better inform ecosystem health, function, and resilience.

**Environmental DNA (eDNA) and nucleic acid-based approaches as a monitoring tool**

Environmental DNA (eDNA) methods focus on the detection and monitoring of sequenced nucleic acid data extracted from environmental samples and can be classed as a form of environmental genomics. They have been successfully used to survey fish biodiversity [22], invasive species [23], detect signature bacteria associated with contaminants [24], assess environmental health and status using indicator organisms [25,26], for species monitoring [23], exploring trophic interactions [27] through to investigating broad biogeographic patterns [28].

Principally, sequence-based eDNA data is a presence/absence or relative abundance measurement used for detection of organisms or function-specific genes (e.g. measuring nitrogen cycle genes in soils to monitor the effects of land-use changes [29]). The use of eDNA data for monitoring has largely focused on taxonomic observations of indicator organisms [30,31]. In the majority of cases, health and function are inferred from the presence/absence or relative abundance of these indicator organisms, with this data then used to inform a response or intervention [32]. Nevertheless, taxonomic observations do not necessarily inform us of the functional performance of an ecosystem. The function can be inferred from taxonomy where we have existing knowledge of an organism’s functional capabilities or through bioinformatic functional inference approaches [33]. However, this is not always straightforward; this is particularly true in the case of microbial indicators. Microbiome metabolic capabilities can be decoupled from taxonomic identity (through gene loss, horizontal gene transfer [34]) and a single organism can exhibit a range of metabolic capabilities depending on environmental conditions [35].
Microbial communities drive key ecosystem processes in natural, managed, and engineered environments [36] and are intrinsically linked to the ecosystem state. Understanding microbiome processes that can mitigate environmental pollutants is therefore critical to maintaining rich biodiversity and healthy ecosystems [37]. Currently, existing and developing knowledge of how microbial communities function naturally and in response to perturbation is not well incorporated into environmental monitoring, surveillance, or management practices [38]. To a large extent, this extends up through producers and consumer taxa — as the interactions and levels of influence of ecosystem connectivity for many taxa are unknown.

Functional profiles of organisms or communities can be produced using DNA-based methods such as metabarcoding of functional genes in biochemical pathways or whole-metagenome sequencing (WMS) which has emerged as a powerful tool to survey biological community structures and their predicted function [39]. Additional analysis that complements and expand traditional metagenomic profiling and targeted eDNA metabarcoding techniques, capturing species-specific and microbial community functional activities are however, still needed [40].

DNA approaches, like those described above, target the genetic or functional potential of organisms or communities, not the realised functional activity or response. Efforts have been made to link gene and transcript abundances of genes associated with specific functions (e.g., ammonia oxidation, denitrification, etc.) with rates of activity, but success has been variable [41,42]. Transcriptomes, sequences of RNA from transcribed genes, take us a step closer to realised function by identifying active organisms and genetic pathways. The presence of specific transcripts does not, however, indicate the associated function is taking place since regulation can occur after expression [43]. Many enzymes also function to catalyse reactions in both directions, making it difficult to know the outcome of activity from gene/expression alone. Therefore, both DNA and RNA approaches are limited in that they do not represent the realised functional components of an environmental response. This is determined by the proteins translated from the RNA, transcribed from the DNA. The metabolome (metabolite content) is then the evidence of the protein activity. Proteomics and metabolomics used in an integrated ecosurveillance framework have the potential to improve our understanding of the realised ecosystem functional state, and with further development, mechanistic models of measurable health and function that can be used to improve management outcomes [23]. However, while DNA and RNA approaches have been readily accessible due to advances in technology for sequencing, quantifying, and comparing these data, until recently, proteomics and metabolomics lacked the depth and sensitivity to be useful except in very targeted experimental approaches. This is true of both more recently targeted organic metabolites (the focus of this review) and those inorganic metabolites we often consider readily measurable (e.g. the intermediates and end products of oxidative and reductive N processes), but which are very difficult to measure at appropriate temporal and spatial scales in situ. With the development of new technology and computational tools, however, applications outside of the laboratory setting are becoming more common.

**Monitoring function (metaproteome and metabolome)**

Recent advances in (meta)proteomics and (community) metabolomics have provided a link between genomic expression and functional characterisation of ecosystem taxa [44]. This provides valuable insight into their in situ metabolism and function, under a range of biotic and abiotic perturbations that genomics alone cannot provide. Table 1 provides an overview of some recent omics-based ecosurveillance applications, which to date, have been predominately microbiome-based.

**Proteomics and metaproteomics applications**

Proteomics has been applied to investigate and monitor the effect of abiotic factors in selected microbiomes, wild species, and model organisms; however, its application to broader ecological monitoring remains underdeveloped. Where a proteomic investigation has been applied to abiotic stress models, much focus has been placed on contaminated microbiomes [67,68] and aquatic species such as fish, crustaceans, and molluscs [69].

As an example, the proteome of goldfish (Carassius auratus) was shown to produce significant changes to oxidative stress and apoptosis inhibition, with no mortality, when subjected to herbicide and fungicide mixtures (8.4 and 42 μg L⁻¹, respectively) and high temperatures (22 and 32°C) [70]. Likewise, the exposure of mature male and female White Sucker fish (Catostomus commersonii) to oil sands-related chemicals in the Athabasca River in Canada showed altered lipid and endocrine metabolism perturbations [71]. The exposure to two different doses of commercial herbicide led to growth impairment and perturbation of the hepatic proteome of rainbow trout (Oncorhynchus mykiss) [72].
### Table 1 Applications of ecosurveillance research with integrated functional datasets intended for omics-based ecosurveillance

| Matrix                      | Objective                                                                 | Omics                                                                 | Reference                  |
|-----------------------------|---------------------------------------------------------------------------|----------------------------------------------------------------------|---------------------------|
| Soil                        | Measure the functional and phylogenetic responses of the microbial community impacted by drought. | Metagenomics: 16S rRNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rRNA gene (bacteria) (Meta) proteomics: LC–MS/MS (Orbitrap) | Bastida et al. [45]       |
| Soil                        | Detect active DNA viruses and RNA viruses in a native prairie soil and determine their responses to extremes in soil moisture | Metagenomics: 16S rRNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rRNA gene (bacteria) (Meta) proteomics: Viral peptides LC–MS/MS | Wu et al. [46]            |
| Soil                        | Assess the metaphenomic responses of a native prairie soil microbiome impacted by drought | Metagenomics: 16S rRNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rRNA gene (bacteria) (Meta) proteomics: GC–MSD (Single quadrupole) | Roy Chowdhury et al. [47] |
| Soil                        | Assess microbial community compositions and functions in response to drought and rainfall events | Metagenomics: 16S rRNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rRNA gene (bacteria) (Meta) proteomics: LC–MS/MS (Orbitrap) | Liu et al. [48]           |
| Soil (microcosm)            | Assessing organic matter decomposition and nutrient cycling in wetland soils | Metagenomics: 16S rRNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rRNA gene (bacteria) (Meta) proteomics: 1H NMR (600 MHz) | McGivern et al. [49]      |
| Soil                        | Assessing contaminants on agricultural microbiome metabolism | Metagenomics: 16S rRNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rRNA gene (bacteria) (after Xu et al. [50]) (Meta) proteomics: LC–MS/MS (Orbitrap) | Chen et al. [51]          |
| River                       | Assessment of surface water quality from multiple non-point source contaminants | Metagenomics: 16S rRNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rRNA gene (bacteria) (Meta) proteomics: GC–MSD (Single quadrupole) | Beale et al. [52]         |
| Soil (-root interface)      | Investigated the symbiotic associations between plant roots with rhizospheric bacterial communities under differing acid mine drainage pollution | Metagenomics: 16S rDNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rDNA gene (bacteria) (Meta) proteomics: LC–TQ–MS | Kalu et al. [53]          |
| Soil                        | Responses of soil microorganisms to polycyclic aromatic hydrocarbon stress | Metagenomics: 16S rDNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rDNA gene (bacteria) (Meta) proteomics: GC–QToF–MS | Li et al. [54]            |
| Sediment/ Water (microcosm) | Response of indigenous microbial structure and functional dynamics in different marine environmental | Metagenomics: 16S rDNA gene (bacteria) • ITS (fungal) (Meta) transcriptomics: 16S rDNA gene (bacteria) (Meta) proteomics: Predicted from PICRUSt | Neethu et al. [55]        |

Continued
| Matrix            | Objective                                                                 | Omics                                                                                     | Reference                  |
|-------------------|---------------------------------------------------------------------------|-------------------------------------------------------------------------------------------|----------------------------|
| Marine Sediment   | Measure the influence of estuarine macrophytes on sediment microbial function and metabolic redundancy | 16S rRNA gene (bacteria) with functional prediction of genes (PICRUSt)                     | Shah et al. [56]           |
| River (flumes)    | Impact of sulfamethoxazole on a riverine microbiome                       | 16S rRNA gene (bacteria)                                                                  | Borsetto et al. [57]       |
| Marine Sediment   | Investigated microbial methane oxidation at the sediment–water interface of a shallow marine methane seep | 16S rRNA gene (bacteria)                                                                  | Taubert et al. [60]        |
| Permafrost        | Reconstruction of fossil and living microorganisms in ancient permafrost  | 16S rDNA gene (bacteria)                                                                  | Liang et al. [61]          |
| Sediment          | Measuring the kinetics of biogeochemical processes in natural and engineered environmental systems | 16S rRNA gene (bacteria and archaea)                                                      | Li et al. [62]             |
| Soil              | Investigated synergistic interactions in a bisphenol A (BPA)-degrading microbial community | 16S rDNA gene (bacteria) with 16S rRNA-tag pyrosequencing                                 | Yu et al. [63]             |
| Sediment (microcosm) | Elucidate the mechanisms driving the rapid biodegradation of Deepwater Horizon Oil in intertidal sediments | 16S rDNA gene (bacteria) with 16S rRNA                                                 | Karthikeyan et al. [64]    |
| Soil              | Investigate soil fungi and their relation to edaphic and environmental variables across three ecosystems | 18S rRNA gene with LC-MS/MS (Orbitrap)                                                   | Fernandes et al. [65]      |

Continued
Continuous exposure to environmental pollutants has been shown to damage the redox response and detoxification processes in crustaceans [73]. Bivalves exposed to metals such as cadmium, copper, lead, and zinc have shown an increased abundance of proteins associated with stress responses, cytoskeletal activity, and protein synthesis [74]. A recent multi-omics study from our group, investigating freshwater turtles exposed to PFAS, identified signs of elevated immune activity and perturbed lipid transport and binding [3]. In addition to monitoring the aquatic environment, transcriptomic- and proteomic-based experiments have provided evidence that herbicides can affect life cycle mechanisms including moulting and the reproduction process of the springtail *Folsomia candida* [75]. Each of these studies presents excellent knowledge development in the assessment of abiotic stress impact on environmental samples or models, but the transition of this knowledge to practical ecosurveillance application remains unfulfilled.

The expansion of metaproteomic approaches that are targeted towards the study of how microbes contribute to ecosystem services [76], capturing both phylogenetic and functional information would help address this. Environmental microbiomes are highly diverse but are currently largely under-represented in public proteomics databases. Furthermore, functional characterisation using metaproteomics is usually performed with the aid of metagenomic sequences acquired for the same sample. Truly representative and diverse metagenomic datasets are difficult to assemble, and therefore, the utility of existing high-quality theoretical proteome databases covering many known isolates eliminating the need for sequencing, is often preferred [77]. A recent proteome-wide study conducted on the taxonomy of life has shown that even after using organism-specific genome and transcriptome resources, ~40% of the identified proteins did not have any functional annotation for their biological processes [78]. Thus, the success of eco-omics-based studies will require improvement in proteome and metaproteome analysis which can reveal the over-representation of functional classes to obtain a global view of environmental health rather than a species-specific view.

Analysis of complex environments is often hindered by the heterogeneity of the sample matrix and its varying concentrations of interfering substances (i.e. salts, humic, fulvic, and tannic acids) [79] that can negatively impact extraction and recovery efficiencies [44,80]. Commercial kits are currently available for the co-extraction of DNA and RNA, but the inclusion of proteins and metabolites requires more research and development [81,82].

**Table 1** Applications of ecosurveillance research with integrated functional datasets intended for omics-based ecosurveillance

| Matrix | Objective | Metagenomics | (Meta) transcriptomics | (Meta) proteomics | Metabolomics | Reference |
|--------|-----------|--------------|------------------------|-----------------|--------------|-----------|
| Sediment | Measure the biological impacts across multiple trophic levels of offshore oil and gas drilling and production operations | 16S rDNA (bacteria) | 16S rRNA (bacteria) | 18S rDNA (eukaryotes) | 18S rRNA (eukaryotes) | Laroche et al. [66] |

Metabolomics and community metabolomics applications

Metabolomics is well suited to assess sublethal biological effects of contaminants and chemical mixtures; it relates chemical processes, intermediates, and end-products of an organism’s metabolism and is closely linked to an exposure-induced phenotype. Coupling quantitative and qualitative chemical analyses with environmental metabolomics bioassays for a range of species/ages/sexes/developmental stages, as is currently underway, will be particularly pertinent in establishing omics-based models to understand the contaminant exposure and impact pathways. For example, metabolomics has recently been applied to zebrafish exposed to environmentally relevant levels of climbazole, a topical antifungal agent, to elucidate the biochemical reasons for reproductive abnormalities seen in female fish [83]. Metabolomics methods have been used to elucidate the sublethal effects of toxicants on a large range of species [84–88].

Where environmental metabolomics comes to the forefront in ecosurveillance applications is with its application to deep data science investigations, coupled with monitoring metadata and other omics datasets for
investigating ecosystem homeostasis [89]. For example, metabolomics has been used to identify correlations between the root microbiome and plant gene translocation in varying plant functions when perturbed by mine drainage pollution that ultimately improved its adaptability and phyto remediation potential [53]. Integrative analyses of transcriptomes and metabolomes in microalgae (Raphidocelis subcapitata) treated with the antibiotic clarithromycin [90] identified impacts to biosynthesis and photosynthesis, highlighting the inhibitory effects of macrolide antibiotics.

Of particular interest is our current understanding of microbial function within environmental microbiomes, which stems from conventional ecology-based surveys and the utility of more recent environmental genomics approaches (eDNA). Attempts to harmonise these data with physicochemical parameters of biotic/abiotic ecosystem (dys)function arising from environmental metabolomics data have been limited and show varying levels of success [9,21,52]. This is potentially biased towards a limited group of microbes (e.g. by the choice of primers), a selection of discrete sampling points, and/or failure to link microbial diversity with functions related to biogeochemical cycles or well-defined metabolic endpoints [48]. So, while resilience and redundancy are theoretically assessed (via, for example, amplicon sequencing), an actual measurable function that is quantifiable is often not. More recent research by Shah et al. [91] employed phylogenetic reconstruction methods to infer genome content and predict functional (relative) abundances that can be matched to metabolite features that are either expressed or consumed within the analysed microbiome.

Like metaproteomic approaches, community metabolomics (e.g. metabolomics of entire communities, usually microbial) may provide functional information post enrichment without the need for 16S rRNA community diversity profiles. Currently, this method has been largely limited to use in soil [92,93] and some specific aquatic systems [52,94]. Just as eDNA can be amplified and used to give an idea of what organisms are present in aquatic systems, so could an environment’s community metabolic profile be preconcentrated, cleaned-up (e.g. via solid-phase extraction), and measured in large aquatic systems such as lakes and streams. These data will generate information on the environmental metabolome and the impact of environmental perturbations on the community. It may also elucidate specific functions such as host-pathogen interactions and plant signalling compounds, and associated responses that are aggregated over physicochemical cycles (i.e. spring-neap tide cycles, etc.) [56,91].

One potential weakness of metabolomics is that results tend to represent the situation at a particular point in time (when the samples were taken). Organisms, however, exist in time and changes are dependent on developmental stage, and external factors, such as climate and health or symbiotic relationship, that can affect an organism’s susceptibility to a pollutant (or pollutants) and thus its potential risk. For it to be a useful tool for monitoring, we need environmental metabolomics to help us understand how differences in timing and duration of exposure (and subsequent deprecation) influence metabolite profiles and organism health. This could be achieved through long-term multi-generational studies, short-term diurnal flux sampling, or by combining existing metabolomics data from model species in the literature and studies performed on the same organism with the same pollutant, but at different life-history stages to give a more comprehensive overview of effects. The integration of in vitro metabolomics with high-throughput screening platforms such as those recently demonstrated by Malinowska et al. [95] may also help with this aim.

**Integrating multi-omic datasets for ecosurveillance**

The expansion of multi-omics research has driven the development of new tools and web-based applications that facilitate their integration for deeper interpretations that extend beyond the correlation of biological molecular features towards biological causality and response. Recent expansion, and a renewed focus on the human gut microbiome, has also led to the development of microbiome-centric tools that pair microbiome sequencing (16S rRNA gene amplicons, shotgun metagenomics, and metatranscriptomics) datasets with analysed metabolomes (mass spectrometry and nuclear magnetic resonance spectroscopy data) [96]. While some of these tools have predominantly been driven by medical and clinical research (e.g. MIMOSA2), they have proven utility when applied to environmental datasets and omics-based ecosurveillance.

Examples of these ‘clinical’ tools being applied within an environmental context are growing. Hua et al. [97] applied MIMOSA2 to investigate the gastrointestinal microbiome of zebrafish (Danio rerio) exposed to the organochlorine pesticide dieldrin, coupling sequence datasets with measure metabolite data. Shah et al. coupled 16S rRNA gene sequencing datasets with untreated metabolomics data to assess the function of marine sediments in tropical estuaries [91] and differing macrophyte zones [56]. Table 2 provides a summary of the recent tools that are freely available; a more expansive list is provided in the review by Pinu et al. [98].
It is now viable to couple ecosurveillance and monitoring data within a multi-omics framework, with computational methods to create novel and integrated ecosystem-scale data frames of system function, organism health, and ecological productivity. This would highlight links between different taxonomic levels, pulling together common metabolic features (harmonisation) into a unified predictive model of ecosystem service provision and system health/trajectory. Such an approach that harmonises conserved metabolic traits amongst taxa with emergent properties of concern (i.e. chemicals and physical attributes), identified via non-target screening, could be utilised in ecosystem predictive models to improve management intervention opportunities, thereby enabling better management decisions and policies.

### Concluding remarks and future perspectives
A critical mass in genomic resources, analytical technologies, and bioinformatics approaches has provided unprecedented insights into the composition, structure, function, and control of the genome, transcriptome, proteome, and metabolome, shedding light upon numerous known and unknown biological pathways and

| Tool                  | Source                                      | Description                                                                 | Data inputs                                                                 | Reference    |
|-----------------------|---------------------------------------------|-----------------------------------------------------------------------------|----------------------------------------------------------------------------|--------------|
| Web of microbes       | http://webofmicrobes.org                   | Web-based exometabolomics data repository and visualization tool.            | • None. Data mining tool.                                                 | Kosina et al. [99] |
| MIMOSA2               | http://elbo-spice.cs.tau.ac.il/shiny/MIMOSA2shiny/ | Web-based and R-based metabolic network tool for inferring mechanism-supported relationships in microbiome-metabolome datasets. | • Taxonomic and/or functional abundances                                 | Noecker et al. [100] |
| MelonnPan             | http://huttenhower.sph.harvard.edu/melonnpan | R-based tool for computational framework modelling to predict community metabolomes from microbial community profiles. | • Metabolite data table (KEGG or HMDB)                                    | Mallick et al. [101] |
| MicrobiomeAnlayst     | https://www.microbiomeanalyst.ca/           | Web-based tool for the comprehensive analysis of common data outputs generated from microbiome studies. Provides a prediction of function based on species annotations. | • Taxonomic and/or functional abundances                                 | Chong et al. [102] |
| Reactome              | https://reactome.org/                       | Web-based multi-omics data visualization and metabolic mapping tool of known biological processes and pathways | • Multi-omics datasets (multiple common formats)                         | Griss et al. [103] |
| PaintOmics 3.0        | http://www.paintomics.org/                 | Web-based tool for the joint visualization of genomics/ transcriptomics, proteomics, and metabolomics data. | • Multi-omics datasets (multiple common formats)                         | Hernández-de-Diego et al. [104] |
| mixOmics              | http://mixomics.org/                       | An R-based multivariate tool that is suited to large ‘omics data sets where the number of variables (e.g. genes, proteins, metabolites) is much larger than the number of samples. | • Transcriptomics, metabolomics, proteomics, microbiome/ metagenomics     | Rohart et al. [105] |
| OmicsAnalyst          | https://www.omicsanalyst.ca/               | Web-based data-driven multi-omics integration tool via intuitive visual analytics | • Transcriptomics, proteomics, metabolomics, and mRNA data                | Zhou et al. [106] |
| OmicsNet 2.0          | https://www.omicsnet.ca/OmicsNet/home.xhtml | Web-based data-driven multi-omics integration tool via Knowledge-based networks | • Transcriptomics, proteomics, metabolomics, and mRNA data                | Zhou and Xia [107] |
phenotypes. Though omics technologies are constantly being adapted to ecological research, there are still limitations and challenges that need to be considered to harness their full potential and acceptance by industry and regulators. In particular, for the application of proteomics to ecological monitoring, the primary issues such as sample and genetic heterogeneity, and limited genomic resources for non-model species complicate data interpretation and limit the potential for integration with other ‘omes’ to obtain systems-level information [108]. Improvement of sampling techniques such as ‘single-pot’ sample extractions could be useful in ensuring the use of a single sample (tube) for multiple omic measurements for a more resolved data interpretation. Additionally, de novo genome sequencing and species-specific database construction would be valuable to identify and validate markers for monitoring purposes. The consideration of intra- and inter-species population diversity, genetic polymorphism, phenotypic plasticity, and developmental stages (including the alternative splicing, post-translational modification) should also be taken into account to inform proteome measurements and other omics-based outputs to better understand taxonomically similar species on a system level [109].

Another area that will likely advance in the future is metabolite identification [110]. At present, many of what is labelled as ‘features’ in metabolomics datasets are not identified [111]. This limits our potential gain in knowledge and understanding of environmental processes. However, collecting robust environmental metadata around these unidentified features could allow them to be correlated and characterised into an environmental context, even if we don’t know their specific function (e.g. metabolite feature X always occurs within Y environments with high metal loads, etc.).

There is currently a paucity of proteome sequence databases available for non-model species. The incompleteness of sequence databases and their limited annotations are often considered a bottleneck for environmental proteomics experiments. Thus, the precise assembly and annotation of genomic and transcriptomic resources would be key to decoding and monitoring proteome level changes in non-model species and their relationship to environmental health. Similar limitations occur in metabolomics. At this point, two main strategies for dealing with metabolomics datasets (which tend to be very large) have involved (1) the establishment of spectral databases to aid with individual feature identification, e.g. the Human Metabolome Database (www.hmdb.ca) [112] and METLIN (https://metlin.scripps.edu/) [113], and (2) developing workflows and analytical packages to facilitate multivariate statistics on individual experimental outcomes. The creation of interactive and open-access databases of pollutants such as the toxic exposome database (http://www.t3db.ca) [112], DrugBank (https://go.drugbank.com) [115], and the EPA’s non-targeted analysis (NTA) database [116] have helped, but each lists data on contaminants/toxicants, not the metabolic response(s) to such compounds. What would help in the future is a library of metabolite profiles for model species exposed to specific pollutants, or mixtures of pollutants as a dedicated tool to facilitate environmental monitoring in complex aquatic environments. The knowledge and infrastructure from existing metabolomics databases could be used for data management, interactive storage, and access to such a system, but it would be reliant on high-quality data from the community to function. Such a database, that is publicly available and easily searchable would facilitate the use of metabolomics in environmental science by allowing scientists to compare results of the analysis of a system, to the metabolic response(s) of the organisms to known pollutants/toxicants (further building upon the ‘Web of microbes’ exometabolomics database for linking chemistry and microbes [99]).

Once a comprehensive database is available, data acquisition techniques such as data-independent acquisition (DIA)-based approaches could be applied to monitor continuous changes. These label-free proteome measurement approaches bring the best of shotgun and targeted acquisition, allowing deep proteome analysis and accurate and reproducible quantitation of proteomes [117], including meta-proteomes [118]. This has recently been supported by advanced machine learning-based tools such as the DIA-NN software [119], which uses neural networks to determine ‘real signals’ from the noise for quantitation and interference removal without any retention time alignment and also makes use of the ProSigt [120] ML predictor to prepare synthetic peptide spectral libraries for quantifying thousands of proteins without a previous observation being required.

Multiple levels of omics-based data coupled within an ecosurveillance approach will bring a greater understanding of the natural and perturbed environment, particularly the microbial systems, upon which we ultimately rely to remediate contaminants. This will be of great benefit to human and environmental health. However, for these multi-omic studies to be conducted in parallel with current approaches to demonstrate their value-add to the status quo, regulators and funding models need to account for the perceived inherent risk of trialling these new approaches (and allow for financial mechanisms that cover the additional analytical costs and help to carry any regulatory risk of just an omics-only approach). Furthermore, agencies must allow for open-ended studies that include (or rely on) nontargeted data but also encompass repeat non-target/omics.
measures over time (i.e. monthly and yearly) that can seem very open-ended to a regulator. It is only then that multi-omics guided ecosurveillance could demonstrate a pathway for improved management intervention opportunities, management decisions, and policies since there are currently limited practical examples of these tools guiding these processes and decisions within the environmental regulatory framework.

In closing, we encourage all readers to explore the opportunities in this exciting area of omics-based research, and as a community of researchers and practitioners, to demonstrate their value and keep pushing these approaches until they become part of the regulatory framework and are embedded in ecosystems monitoring programs.

## Summary

- Current environmental monitoring efforts often focus on known, regulated contaminants ignoring the effects of unmeasured compounds, environmental factors, and subtle biochemical perturbations.

- Metabolomics- and proteomics-based approaches can be coupled with DNA/RNA sequence technologies to provide measured functional outputs.

- Using multiple levels of omics technology-based assessments together into an ecosurveillance approach will bring a greater understanding of the natural and perturbed environment with great benefit to environmental health.

## Competing Interests

The authors declare that there are no competing interests associated with the manuscript.

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## Abbreviations

DIA, data-independent acquisition; EDA, effects directed analysis; PFAS, perfluoroalkyl substances.

## References

1. ANZECC and ARMCANZ. (2018) Australian and New Zealand Guidelines for Fresh and Marine Water Qualit. [cited 2021; Available from: https://www.waterquality.gov.au/anz-guidelines

2. Ebner, J.N. (2021) Trends in the application of “omics” to ecotoxicology and stress ecology. *Genes* **12**, 1481 https://doi.org/10.3390/genes12101481

3. Beale, D.J., Hillyer, K., Nilsson, S., Limpus, D., Bose, U., Broadbent, J.A. et al. (2022) Bioaccumulation and metabolic response of PFAS mixtures in wild-caught freshwater turtles (Emydura macquariimacquarii) using omics-based ecosurveillance techniques. *Sci. Total Environ.* **806**, 151264 https://doi.org/10.1016/j.scitotenv.2021.151264

4. Beale, D.J., Nilsson, S., Bose, U., Bourne, N., Stockwell, S., Broadbent, J.A. et al. (2022) Bioaccumulation and impact of maternal PFAS offloading on egg biochemistry from wild-caught freshwater turtles (Emydura macquarii macquarii). *Sci. Total Environ.* **817**, 153019 https://doi.org/10.1016/j.scitotenv.2022.153019

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94 Beale, D.J., Barratt, R., Marlow, D.R., Dunn, M.S., Palombo, E.A., Morrison, P.D. et al. (2013) Application of metabolomics to understanding biofilms in water distribution systems: a pilot study. *Biofouling* **29**, 283–294 https://doi.org/10.1080/08927014.2013.772140

95 Malinowska, J.M., Paksaat, T., Sund, J., Carpi, D., Boughid, M., Weber, R.J.M. et al. (2022) Integrating in vitro metabolomics with a 96-well high-throughput screening platform. *Metabolites* **18**, 11 https://doi.org/10.3390/metabo9040076

96 Yin, X., Altman, T., Rutherford, E., West, K.A., Wu, Y., Choi, J. et al. (2020) A comparative evaluation of tools to predict metabolite profiles from microbiome sequencing data. *Front. Microbiol.* **11**, 595910 https://doi.org/10.3389/fmicb.2020.595910

97 Hua, Q., Adamovsky, O., Veselcová, H., Boyda, J., Schmidt, J.T., Kozuch, M. et al. (2021) Microbiome analysis and predicted relative metabolomic turnover suggest bacterial heme and selenium metabolism are altered in the gastrointestinal system of zebrafish (Danio rerio) exposed to the organochlorine dieldrin. *Environ. Pollut.* **268**, 115715 https://doi.org/10.1016/j.envpol.2020.115715

98 Pinar, F.R., Beale, D.J., Paton, A.M., Kourmenos, K., Swarm, S., Schirra, H.J. et al. (2019) Systems biology and multi-Omics integration: viewpoints from the metabolomics research community. *Metabolites* **9**, 76 https://doi.org/10.3390/metabo9040076

99 Kosina, S.M., Greiner, A.M., Lau, R.K., Jenkins, S., Banar, R., Bowen, B.P. et al. (2018) Web of microbes (WoM): a curated microbial exometabolomics database for linking chemistry and microbes. *BMC Microbiol.* **18**, 115 https://doi.org/10.1186/s12866-018-1256-y

100 Noecker, C., Eng, A., Muller, E. and Borenstein, E. (2022) MIMOSA2: a metabolite network-based tool for inferring mechanism-supported relationships in microbiome-metabolite data. *Bioinformatics* **38**, 1615–1623 https://doi.org/10.1093/bioinformatics/btat003

101 Mallick, H., Franzosa, E.A., McVeer, L.J., Banerjee, S., Sirota-Madi, A., Kostic, A.D. et al. (2019) Predictive metabolomic profiling of microbial communities using amplicon or metagenomic sequences. *Nat. Commun.* **10**, 3136 https://doi.org/10.1038/s41467-019-10927-1

102 Chong, J., Liu, P., Zhou, G. and Xia, J. (2020) Using microbiomeAnalyst for comprehensive statistical, functional, and meta-analysis of microbiome data. *Nat. Protoc.* **15**, 799–821 https://doi.org/10.1038/s41596-019-0264-1

103 Griss, J., Viteri, G., Sidiropoulos, K., Nguyen, V., Fabregat, A. and Hermjakob, H. (2020) ReactomeGSA - efficient multi-Omics comparative pathway analysis. *Mol. Cell. Proteomics* **19**, 2115–2125 https://doi.org/10.1074/mcp.M120.002155

104 Hernández-de-Diego, R., Tarazona, S., Martínez-Mira, C., Balzano-Nogueira, L., Furió-Tarí, P., Pappas, Jr, G.J. et al. (2018) Paintomics 3: a web resource for the pathway analysis and visualization of multi-omics data. *Nucleic Acids Res.* **46**, W503–W509 https://doi.org/10.1093/nar/gky466

105 Rohart, F., Gautier, B., Singh, A. and Lê Cao, K.-A. (2017) Mixomics: an R package for analysis of multi-Omics data. *PLoS Comput. Biol.* **15**, e1005752 https://doi.org/10.1371/journal.pcbi.1005752

106 Demichev, V., Messner, C.B., Vernardis, S.I., Lilley, K.S. and Ralser, M. (2020) DIA-NN: neural networks and interference correction enable deep proteome coverage in high throughput. *Nat. Commun.* **11**, 2115 https://doi.org/10.1038/s41592-020-0942-5

107 Zhou, G. and Xia, J. (2019) Using omicsNet for network integration and 3D visualization. *Bioinformatics* **35**, es300550 https://doi.org/10.1093/bioinformatics/btx765

108 Mancia, A. (2018) New technologies for monitoring marine mammal health. *Mar. Mamm. Ecol. Med.* **29**, 251–260 https://doi.org/10.1007/s12265-018-0143-x

109 Jubeaux, G., Audouard-Combe, F., Simon, R., Tutundjian, R., Salvador, A., Geffard, O. et al. (2012) Vitellogenin-like proteins among invertebrate species diversity: potential of proteomic mass spectrometry for biomarker development. *Environ. Sci. Technol.* **46**, 6315–6323 https://doi.org/10.1021/es305505h

110 Dias, D.A., Jones, O.A.H., Beale, D.J., Boughton, B.A., Benheim, D., Kourmenos, K.A. et al. (2016) Current and future perspectives on the structural identification of small molecules in biological systems. *Metabolites* **6**, 46 https://doi.org/10.3390/metabo6040046

111 Jones, O.A.H. (2018) Illuminating the dark metabolome to advance the molecular characterisation of biological systems. *Metabolites* **14**, 1 101 https://doi.org/10.3390/metabo14010101

112 Wishart, D.S., Feunang, Y.D., Manca, A., Guo, A.C., Liang, K., Vázquez-Fresco, R. et al. (2017) HMDB 4.0: the human metabolome database for 2018. *Nucleic Acids Res.* **46**, D608–D617 https://doi.org/10.1093/nar/gkw1089

113 Xue, J., Guillas, C., Benton, H.P., Warth, B. and Szudak, G. (2020) METLIN MS2 molecular standards database: a broad chemical and biological resource. *Nat. Methods* **17**, 953–954 https://doi.org/10.1038/s41592-020-0942-5

114 Pina, R., Ron, A., Sajed, T., Guo, A.C., Djumoubou, Y. et al. (2015) T3DB: the toxic exposome database. *Nucleic Acids Res.* **43**, D329–D334 https://doi.org/10.1093/nar/gku014

115 Wishart, D.S., Knox, C., Guo, A.C., Cheng, D., Shrivastava, S., Turr, D. et al. (2007) Drugbank: a knowledgebase for drugs, drug actions and drug targets. *Nucleic Acids Res.* **36**, D901–D906 https://doi.org/10.1093/nar/gkm958

116 Ullrich, E.M., Sokus, J.R., Grueke, C.M., Richard, A.M., Newton, S.R., Snyman, M.J. et al. (2019) EPA’s non-targeted analysis collaborative trial (ENTACT): genesis, design, and initial findings. *Anal. Bioanal. Chem.* **411**, 853–866 https://doi.org/10.1007/s00216-018-1435-6

117 Gillet, L.C., Navarro, P., Tate, S., Röst, H., Selevsek, N., Reiter, L. et al. (2012) Targeted data extraction of the MS/MS spectra generated by data-independent acquisition: a new concept for consistent and accurate proteome analysis”. *Mol. Cell. Proteomics* **11**, 0111.016717 https://doi.org/10.1074/mcp.O111.016717

118 Long, S., Yang, Y., Shen, C., Wang, Y., Deng, A., Qin, Q. et al. (2020) Metaproteomics characterizes human gut microbiome function in colorectal cancer. *NPJ Biofilms Microbiomes* **6**, 14 https://doi.org/10.1038/s41522-020-0123-4

119 Demichev, V., Messner, C.B., Vernardis, S.I., Lilley, K.S. and Ralser, M. (2020) DIA-NN: neural networks and interference correction enable deep proteome coverage in high throughput. *Nat. Methods* **17**, 41–44 https://doi.org/10.1038/s41592-019-0638-x

120 Gessualt, S., Schmidt, T., Zoigl, D.P., Samaras, P., Schnattbaum, K., Zerweck, J. et al. (2019) ProSil: proteome-wide prediction of peptide tandem mass spectra by deep learning. *Nat. Methods* **16**, 509–518 https://doi.org/10.1038/s41592-019-0426-7