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State-of-the-art methods for exposure-health studies: Results from the exposome data challenge event



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ABSTRACT

The exposome recognizes that individuals are exposed simultaneously to a multitude of different environmental factors and takes a holistic approach to the discovery of etiological factors for disease. However, challenges arise when trying to quantify the health effects of complex exposure mixtures. Analytical challenges include dealing with high dimensionality, studying the combined effects of these exposures and their interactions, integrating causal pathways, and integrating high-throughput omics layers. To tackle these challenges, the Barcelona Institute for Global Health (ISGlobal) held a data challenge event open to researchers from all over the world and from all expertises. Analysts had a chance to compete and apply state-of-the-art methods on a common partially simulated exposome dataset (based on real case data from the HELIX project) with multiple correlated exposure variables ($P > 100$ exposure variables) arising from general and personal environments at different time points, biological molecular data (multi-omics: DNA methylation, gene expression, proteins, metabolomics) and multiple clinical phenotypes in 1301 mother-child pairs. Most of the methods presented included feature selection or feature reduction to deal with the high dimensionality of the exposome dataset. Several approaches explicitly searched for combined effects of exposures and/or their interactions using linear index models or response surface methods, including Bayesian methods. Other methods dealt with the multi-omics dataset in mediation analyses using multiple-step approaches. Here we discuss features of the statistical models used and provide the data and codes used, so that analysts have examples of implementation and can learn how to use these methods. Overall, the exposome data challenge presented a unique opportunity for researchers from different disciplines to create and share state-of-the-art analytical methods, setting a new standard for open science in the exposome and environmental health field.

1. Background

The exposome is a concept of growing interest in the field of environmental and molecular epidemiology. Described as “the totality of

human environmental exposures from conception onwards”, it recognizes that individuals are exposed simultaneously to a multitude of environmental factors and takes a holistic approach to the discovery of etiological factors for disease. The exposome’s main advantage over

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traditional ‘one-exposure-one-disease’ study approaches is that it provides an unprecedented conceptual framework for the study of multiple environmental hazards (urban, chemical, lifestyle, social) and their combined effects. Indeed, classical single pollutant models make unsure the fact that the analysed association is due to the pollutant effect or from another correlated exposure not taken into account in the analysis. They are also unable to capture interactions and cumulative effects from the exposure mixture. Furthermore, given the increasing availability of complex environmental health data due to the emergence of new technologies (such as electronic health records, high throughput omics platforms, wearable sensors, etc), there is a need for more advanced statistical approaches that focus on complex mixtures of exposures.

However, the analysis of such complex data comes with numerous challenges, for instance the usually high correlations between exposures of the same family (air pollutants, lifestyle), the ability to capture cumulative low dose effects, assess interactions, identify important components of the mixture. Recently, methods have been applied to take into account multiple exposures and the interactions between them, for example by using mixture analysis methods, by integrating the selection, shrinkage and grouping of correlated variables (e.g. LASSO, elastic-net, adaptive elastic-net), dimension reduction techniques (e.g. principal component, partial least square analysis) or bayesian model averaging (BMA), Bayesian kernel machine regression (BKMR), etc.) (Stafoggia et al., 2017; Lazarevic et al., 2019). A series of limitations of these approaches have been previously identified such as the lack of model selection stability (shrinkage methods), lack of interpretability of the latent variables (dimension reduction) and computational inefficiency (Bayesian models). In addition, they are rarely applied in the context of large (>100 variables) and heterogenous exposome data (omics, categorical/continuous variables).

To address the numerous challenges that come with the analysis of newly available exposome data and to promote interdisciplinary collaboration between researchers from around the world, ISGlobal hosted a 3-day online data challenge in April 2021 entitled “the Exposome Data Challenge Event”. This is the first data challenge organised in this field, and still a rarity in the academic sphere. Data challenges, hackathons or crowdsourcing events were initially used for software development in the 2000’s but are now additionally used in healthcare research to accelerate innovation and peer-reviewing, structure learning, test reproducibility of results, and enable wide participation. Briefly, these events allow participants, usually organized in teams, to respond in a short time frame (1–3 months) to common biological questions using a specifically provided dataset. The Exposome Data Challenge Event was inspired by previous events such as the National Institute of Environmental Health Sciences (NIEHS) workshop for assessing Health Effects of Environmental Chemical Mixtures in Epidemiology (Taylor et al., 2016) organised in 2015 or the DREAM challenge annual series (Ellrott et al., 2019) during which biological data sets are released to the international community to build computational models that address specific biological questions. This data challenge was particularly motivated by the need to address interpersonal interactions constrained by the COVID-19 pandemic and provide a platform to researchers from various genders, backgrounds, and career stages. It was built upon a well-established dataset from the HELIX cohorts which measured the early life exposome (Maitre et al., 2018; Vrijheid et al., 2014). Exposome-health and exposome-omics associations in this dataset have been previously well described within the HELIX consortium. However, the consortium was ready to open and brainstorm with a wider scientific community about unresolved challenges of exposome cohort data such as non-linear combined effects of exposures, causality and omics integration, repeated time points and multi-cohort design. Our main objective was to promote innovative statistical, data science, or other quantitative approaches to study the health effects of complex multi-dimensional exposures and high throughput omics measurements in this unique exposome dataset. At this stage, exposome studies are often framed without an a priori hypothesis, with an

explorative approach, therefore we formulated the challenges around the data analysis and not specific research questions, ensuring the generalizability of the outputs. In addition, we wanted to deliver for the community a common public training dataset for exposome studies, programming code clearinghouse and open collaborations for future projects.

This event gathered a widely diverse scientific audience of 307 participants, including environmental epidemiologists, biostatisticians and computational scientists, to discuss state-of-the-art statistical methods for studying exposome-health associations. The participants were offered an opportunity to test their statistical methods of choice addressing one or several key challenges: a) the high dimensionality of the data, b) combined effect of exposure or mixtures, c) the omics data integration and the d) causal structure in the exposome. Participants were encouraged to accommodate in their approaches some of the particularities of the data (e.g. multi-cohort, count responses, categorical and continuous exposure variables, exposures measured at two time points, etc). Visualization of the results was also a key point across all the challenges listed above. This report outlines the approaches presented at the event, which represent useful computational, conceptual, and statistical models for analyzing high dimensional exposome datasets, including omics and health outcome associations. In collaboration with the event committee and the selected participants, we discuss the different techniques.

2. Methods

2.1. Event organisation

First, participants were invited to submit an abstract describing their team, the challenge(s) and the method they would apply on a common partially simulated exposome dataset (based on real case data from the HELIX project). The planning committee selected a total of 25 abstracts out of 39 based on method clarity, novelty, relevance for the exposome field and challenges presented. Second, the selected participants were invited to apply their method on the dataset during a month leading to the event. Third, they presented at the event their method’s statistical background, type of research question(s) it best addressed, and their results. At the end of the event, the committee and the audience voted for the best presentations based on clarity, novelty and relevance. Finally, the participants made their code available on the github account (Gonzalez, 2021) of the event and videos of the presentation are available on the [youtube channel of the ATHLETE project](#).

2.2. Data

The exposome data provided for this challenge came from the HELIX subcohort database and were partially simulated. The HELIX study (Maitre et al., 2018; Vrijheid et al., 2014) represents a collaborative project across six established and ongoing longitudinal population-based birth cohort studies in six European countries (France, Greece, Lithuania, Norway, Spain, and the United Kingdom). From the 31 472 mother–child pairs included in the cohorts, a subcohort of 1301 mother–child pairs were followed up with measurements of biomarkers, omics signatures and health outcomes at 6–11 years of age. The data provided for the challenge came from this subsample, but it was partially simulated to respond to policies of data anonymization for privacy protection in the cohorts. In detail, for the set of health outcomes and exposures, we conducted the following process: for each participant, a total of 50 random variables (different for each subject) were converted into missing values and then imputed (in successive rounds of 10 variables at a time) with the method of chained equations. Thus, for each participant, some of the values in the provided dataset were real and some were simulated, in a way that precluded knowing what is real data and what is simulated. Omics data instead were kept intact but annotations of genes and metabolites were not provided. We provided an

imputed dataset in which all missing values in the original data were imputed by the chained equations method. Exposure data were transformed (e.g. logarithmic, square root, tertiles) to achieve symmetric distributions with a homogenous range of values (Annex 1). The original raw data are available on request subject to ethical and legislative review. The “HELIX Data External Data Request Procedures” are available with the data inventory in this website: <https://www.projecthelix.eu/data-inventory>.

The datasets are available in the [github repository](#) of the challenge event and transcriptomic and Epigenomics through a FigShare account: <https://figshare.com/account/home#/projects/98813>. An overview of available data is shown in Fig. 1 and the complete codebook description is available in Annex 1. It includes more than two hundred environmental exposure variables, 13 covariates, six health outcomes [body mass index (BMI), asthma, birth weight, neurobehaviour, intelligence quotient (IQ)], and omics data (serum metabolome, urine metabolome, proteins, gene expression, methylation). Exposure, covariate and health outcome datasets contain both continuous and categorical values; health outcome data additionally included count data. Exposure and omic datasets both included highly correlated features (correlation > 0.8).

2.3. Challenges

The main challenges that were addressed during the workshop were as follows:

- (1) The high dimensionality of exposure data and, more precisely, methods to reduce it while minimizing the loss of useful information regarding health associations and combined effects. The exposome dataset available during the challenge consisted of a large number of environmental exposure variables and a multi-omics dataset (approximately 0.5 M features), with a high inter-variable correlation, but a small sample size ($N = 1301$), typical of exposome studies.
- (2) The combined effect of exposure or mixtures. Researchers are interested in studying individual and combined effects of a large number of exposures together accounting for their potential interactions. Effects of environmental exposures may be small when taken individually, but their aggregation may lead to a

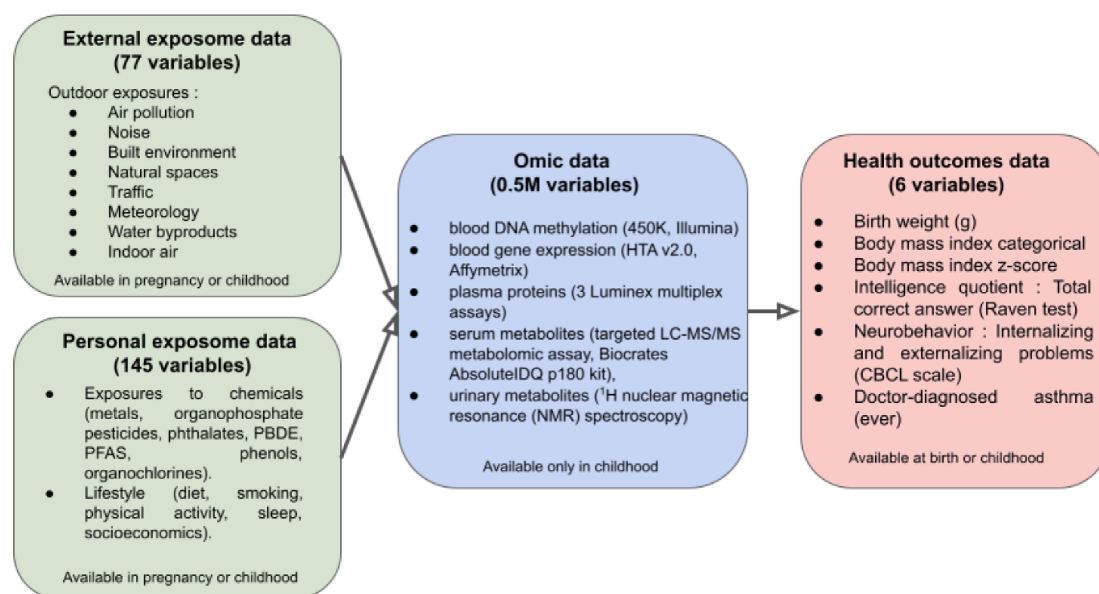
significant alteration of the health outcome of interest, leading to cocktail effects that researchers want to investigate.

- (3) Omics data integration. Omics data may be used, in addition to exposure data, in order to provide causal inference on the link between exposome and health. The challenge here was to incorporate one or several of the different omic layers available, with the purpose of finding patterns that can explain variations in one or more health outcomes and analysing how the exposome and the omes interact with regard to these outcomes. The method used in this context must be able to maximise omics data predictive power with very high dimensional data and small sample size ($N \ll P$).
- (4) Causal structures in exposure data. This challenge included: 1) how to incorporate *a priori* hypothesized causal relationships between the different exposures and one health outcome into the analysis, 2) the comparison of this *a priori* approach with *agnostic* analyses that would perform variable selection treating all exposures in the same way, 3) how one can answer a large number of causal questions referring to different exposures using causal inference techniques for high-dimensional data, 4) the incorporation of mediation analysis and high-dimensional mediation analysis.

Additional points of interest included visualisation techniques, the handling of the multicenter design of the study, the control for potential confounders that may have an effect on the health outcomes and need to be considered when studying associations with the exposomic features, and missing data in exposome datasets.

3. Results

In this section, we summarise the statistical methods used by the participants in the Exposome Data Challenge listed in Table 1. All the codes developed by the participants to perform their analyses are available in the [GitHub repository](#) of the event. Briefly, most presentations focused on one health outcome out of the six available, mainly child BMI, and five used multiple outcomes. Seven presentations included categorical outcomes (*i.e.* asthma, BMI and birth weight < 2500gr), the rest focused on continuous outcomes only. Less than half of the methods included categorical exposure variables, focusing mainly



We also included 13 covariates (6 on mother, 7 on child).

Fig. 1. Overview of the data available during the challenge. The data were based on the HELIX project which collected exposome, omics and health data from six mother-child cohorts across Europe in 1301 participants (Maitre et al., 2018).

Table 1

Summary of all the presentations during the exposome data challenge. Ge = Gene Expression, Me = DNA Methylation, Pr = Proteins, SM = Serum Metabolites, UM = Urine Metabolites, IQ = total correct answer (RAVEN test), Neurobehavior = Internalizing and externalising problems (CBCL scale).

| Presentation order | Authors names, University | Presentation Title | Method names | Link to slides | Link to papers | Omics data | Categorical exposures | Selected health outcome |
|---|---|---|---|----------------------|---|------------|-----------------------|---|
| 2. Approaches to study the combined effect of exposures on health | | | | | | | | |
| 2.1 Approaches explicitly searching for combined effects of exposures, linear or nonlinear, and/or their interactions | | | | | | | | |
| 4 | Chris Gennings, Icahn School of Medicine at Mount Sinai | Evaluating a Mixture Effect of Perinatal Environmental Exposures on Childhood BMI Using Weighted Quantile Sum (WQS) Regression | Weighted Quantile Sum Regression (WQS) | Link | (Carrico et al., 2015) | | N | BMI z-score |
| 6 | Matthew Carli and David Wheeler, Virginia Commonwealth University | Exposome Analysis with Bayesian Group Index Regression | Bayesian Group Index Regression | Link | (Wheeler et al., 2021) | SM | N | Asthma |
| 2 | Vishal Midya, Icahn School of Medicine at Mount Sinai | A novel penalized LASSO type Bayesian Weighted Quantile Sum Regression Approach for Exposome-outcome effect estimation | Bayesian Weighted Quantile Sum Regression | Link | (Colicino et al., 2020) | | N | BMI z-score |
| 5 | Shounak Chattopadhyay, Duke University | Synergistic Interaction Detection | Synergistic Interaction Detection | Link | <i>Article in preparation</i> | | N | Birth weight |
| 3 | Ander Wilson, Colorado State University, Daniel Mork | Exposome Health Association Studies Using Bayesian Treed Distributed Lag Mixture Models | Treed Distributed Lag Mixture Models | Link | (Mork and Wilson, 2021) | | N but could | BMI z-score |
| 22 | Michele Peruzzi, Duke University | Multi-Outcome Meshed Gaussian Processes on Projected Inputs for Scalable Inference with Exposome Data | Meshed Gaussian Processes | Link | (Peruzzi et al., 2020) | | Y | Birth weight, BMI and related variables |
| 1 | Glen McGee, University of Waterloo | Quantifying Exposome-Health Associations with Bayesian Multiple Index Models | Bayesian Multiple Index Models | Link | (McGee et al., 2021) | | N | BMI z-score |
| 2.2 Approaches using Machine Learning to maximise prediction performance | | | | | | | | |
| 14 | Jean-Baptiste Guimbaud, Remy Cazabet, Léa Maitre, LIRIS-ISGlobal | Leveraging machine learning and explainable AI to better understand exposomic data | Multilayer Perceptron, xgboost, random forest, SVM, Elastic-net, SHAP | Link | NA | | Y | All |
| 13 | Fei Zou, University of North Carolina, Chapel Hill | Deep-Exposome: A Predictive and Interpretative Deep Neural Network Ensemble for Exposome Data | Improved Bootstrap Aggregating and PermFIT | Link | (Mi et al., 2021, n.d.) | | Y | All |
| 16 | Lucile Broséus and Paulina Jedynak, Université Grenoble Alpes | Searching for the risk factors for childhood overweight - A novel approach to identify the most relevant child BMI-associated exposures | Univariate Ordinal Logistic Regression and Multiple Correspondence Analysis (MCA) | Link | NA | | Y | BMI categorical |
| 24 | Hua Yun Chen, University of Illinois at Chicago | Estimating the effects of exposome and their interactions | Explained variation (EV) in linear models | Link | NA | | Y | Birth weight, IQ |
| 2.3 Multi-stage approaches for combined effects | | | | | | | | |
| 12 | John Pearce, Medical University of South Carolina | Exposure Continuum Mapping for predicting health and disease in exposome studies | Exposure Continuum Mapping and Generalized Additive Models | Link | (Pearce et al., 2021) | | N but could | Birth weight |
| 8 | Jaime Benavides and Lawrence Chillrud, Columbia University | Pre- and postnatal urban exposure patterns and childhood neurobehavior | Principal Component Pursuit (PCP), Factor Analysis, GAM and LASSO | Link | (Gibson et al., 2021) | | N | Neurobehavior |
| 11 | Sejal Mistry and Ramkiran Gouripeddi, University of Utah | Clustering Exposure Trajectories to Classify Phenotypic Characteristics | clustering transitions on phenotypic characteristics | Link | NA | | N | BMI z-score |
| 23 | Sanjib Basu, Ruizhe Chen, Yu-Che Chung, | Missingness pattern and exposure selection for | CORrelation LearNING and exposure Selection | Link | NA | | Y | BMI z-score |

(continued on next page)

Table 1 (continued)

| Presentation order | Authors names, University | Presentation Title | Method names | Link to slides | Link to papers | Omics data | Categorical exposures | Selected health outcome |
|--|---|--|---|----------------------|---------------------|----------------|-----------------------|--|
| | Jiyeong Jang Mary Turyk and Hua-Yun Chen, University of Illinois at Chicago | mixed-type exposome data | (COLRNS) and A Test for Realized Missing Completely At Random | | | | | |
| 3. Studies using omics data to improve inference on the link between exposome and health | | | | | | | | |
| 10 | Ziyue Wang, National Institute of Environmental Health Sciences | Integrative Analysis and Visualization of Exposome and Transcriptome data | Differential expressed gene analysis (DEG) and Mediation analysis | Link | NA | GE | Y | Asthma |
| 18 | Xiaotao Shen, Stanford University | Decoding unknown links between the exposome and health outcomes by multi-omics analysis | bi-directional mediation analysis. | Link | NA | GE, UM, SM, Pr | N | IQ, Neurobehavior, BMI z-score |
| 19 | Congrong Wang, Brigitte Reimann, Rossella Alfano, Hasselt University | Meet-in-the-middle meets multi-omics: a strategy to identify molecular signatures of environmental drivers of childhood BMI | Multi-omics Mediation Analysis | Link | NA | All | Y | BMI z-score |
| 21 | Miao Yu, Icahn School of Medicine at Mount Sinai | Molecular Gatekeepers bridge the exposome and health | Molecular gatekeepers discovery | Link | (Yu et al., 2021) | SM | N | Asthma |
| 17 | Carl Brunius, Chalmers University of Technology | Omics Modules for Exposome-Health Associations (OMEXA) | MUVR, Generalized Linear Models and Triplot | Link | (Shi et al., 2019) | All | Y | All |
| 9 | Nikos Stratakis, University of Southern California | Latent unknown clustering integrating multi-omics data (LUCID) with phenotypic traits | Unknown Clustering (LUCID) | Link | (Peng et al., 2020) | SM, UM | N | BMI z-score |
| 7 | Qiong Wu, University of Maryland, College Park | A new statistical graph model to systematically study associations between multivariate exposome data and multivariate metabolomics data | Bipartite Graph | Link | NA | SM | N | None |
| 4. Causal inference | | | | | | | | |
| 25 | Charlie Roscoe, Hari Iyer, Huichu Li, and Marcia Pescador Jimenez, Harvard University | Air pollution and childhood cognition: a g-computation approach to assess mediation by a mixture of metals | Causal mediation analysis and quantile g-computation | Link | (Keil et al., 2020) | | N | IQ |
| 20 | Daniela Zugna, Chiara Moccia, University of Turin | Application of a novel method for mediation analysis in the exposome context | Mediation Analysis | Link | (Loh et al., 2020) | | Y | Birth weight (<2500gr) |
| 15 | Alejandro Caceres, ISGlobal | Using causal random forest to determine exposure environments with high sexual dimorphisms | Causal random forests | Link | NA | Me, GE | Y | BMI z-score difference in boys and girls |

Table 2

Method classification according to dimensionality reduction technique used in the exposome data challenge. The number of the presentation corresponds to the list in Table 1. The ones with an * correspond to presentations which can fit in several categories.

| Feature selection | | | | |
|---|------------------------|-------------------------------------|---|---|
| Feature extraction | None | By statistical tests (correlation*) | Regularized regression (LASSO, Elastic-net) | Feature importance (tree based, permutation based, regression coefficients etc..) |
| None | 12, 13, 15, 18, 24, 25 | 7, 21 | 3, 5, 20 | 14, 16, 17 |
| Indices (weighted quantiles, risk scores) | | | 1*, 2*, 10 | 1*, 2*, 4 |
| Feature projection (PCA/FA/PCP/MCA) | 11*, 22 | | 8 | 19 |
| Clusters | 9, 11*, 12, | 23 | | |

Abbreviations: FA, Factor Analysis; PCA, Principal Component Analysis; PCP, Principal Component Pursuit; MCA, Multiple Correspondence Analysis.

*Both correlations with the outcome and within features were used to perform feature selection.

on continuous chemical exposures. Eight presentations included omics data, including one with all omics layers. A separate results section is dedicated to the comparison of the findings across methods that focused on child BMI and chemical exposures.

3.1. Approaches to deal with high dimensionality of the data

Most analyses presented at the event, even those dealing only with environmental exposures ($n > p$), applied some sort of dimensionality reduction techniques, as summarized in Table 2. These techniques can be split in two categories: 1) feature extraction techniques (Khalid et al., 2014), which consist in computing derived variables that are functions of the original ones and have a smaller dimension, but retain/extract most of the information contained in the original feature space, and 2) feature selection techniques which consist in selecting a subset of the original variable set, while keeping most of the information contained in the whole variable space.

Among the feature extraction techniques that were used during the event, we can further define different types: 1) techniques that calculate summary measures (indices) based on the creation of weighted combinations of exposures that predict the health outcome; 2) feature projection techniques such as Principal Component Analysis (PCA) (Jolliffe, 1986), Factor Analysis (FA), Principal Component Pursuit (PCP) (Candes et al., 2009) or Multiple Correspondence Analysis (MCA) (Blasius and Greenacre, 2006) to represent the data in a low dimensional space; and 3) techniques that provide clustering of participants sharing a similar exposome profile, which could be predictive of the outcome, *i.e.* supervised, or not.

Feature selection approaches used during the event can also be divided into different types of selections: 1) based on correlation with the outcome of interest, using Pearson's correlation as a screening approach; 2) based on regularisation (LASSO, Elastic-net regression) by shrinking the less relevant features' coefficients; 3) based on feature importance which reflects the impact of a given feature on the model predictions through permutation (Altmann et al., 2010), random forest (Breiman, 2001), or regression coefficients. Other statistical tests exist for features' selection (ex: chi-squared test, ANOVA, etc..) but were not used during this challenge. Some analysts made an *a priori* selection, by choosing to focus only on a particular subset of exposures, *e.g.* lifestyle exposure, based on prior knowledge for causal models (25) or model abilities (continuous-only-exposures).

Eight studies implemented feature selection but not feature extraction, four implemented feature extraction but not feature selection, and seven applied both techniques (Table 2).

4. Approaches to study the combined effect of exposures on health

4.1. Approaches explicitly searching for combined effects of exposures, linear or nonlinear, and/or their interactions

Several methods were presented to capture the effect of exposure mixtures. Most of them can be classified as linear index models or response surface methods. Linear index models generate new variables (usually called indices) that are weighted averages of the original exposures, and regress those indices against the health outcome. Response surface methods fit a complex high-dimensional surface to the data, and are thus able to capture complex non-linearities and interactions.

In the group of index models, Gennings (4) et al. presented the weighted quantile sum regression (WQSR) method (Carrico et al., 2015), its history and recent extensions. WQSR builds a new index, which is a weighted average of the initial exposures (previously categorised into quantiles as a way to standardise the data and prevent the effect of influential observations). The new index is regressed against a health outcome, producing a single regression coefficient. The weights to build the index, which incorporate directionality constraints (*e.g.*, all

variables are expected to produce negative effects on the health outcome), are estimated simultaneously with the regression coefficients. This technique assumes additive effects of the different pollutants. Some presented extensions included models to produce strata-specific weights and regression coefficients, combining two indices (one for each directionality) in the same model, or using resampling to improve the properties of the method.

Carli (6) et al. presented the application of Bayesian Group Index Regression (BGIR), a Bayesian equivalent to WQSR that does not use directionality constraints and allows multiple indices (based on groups of exposures) in the same model (Wheeler et al., 2021). In the application, they included several indices according to exposure families, and each exposure family could have two indices if the family contained both exposures that were positively and negatively correlated with the outcome. They conducted analyses separately by cohort and included serum metabolomics data as an additional group of exposures. Midya (2) et al. presented the application of LASSO-type Bayesian Weighted Quantile Sum Regression (LBWQSR), which is similar to BGIR, but introducing LASSO and Elastic net penalties to prevent overfitting (Xu and Ghosh, 2015).

In the group of response surface methods, Chattopadhyay (5) et al. presented a method to search for two-way non-linear interactions. Two-dimensional splines were used to capture the shape of the association for all pairs of continuous exposures. A Bayesian paradigm was used, with priors that allow shrinking terms to zero in the absence of interaction. Prior information on the direction of the interaction (synergistic vs. antagonistic) was also incorporated.

Mork et al. (3) presented the application of Treed Distributed Lag Mixture Models (TDLMM) (Mork and Wilson, 2021). This method uses a Bayesian additive regression trees style model that performs exposure selection of main effects and two-way interactions, and incorporates the repeated exposure measurements available at two time points. The model performs hierarchical variable selection (interactions are only included if both main effects are included), performs shrinkage of regression coefficients, and performs dimension reductions by averaging over multiple time points when there is no evidence that the association varies over time.

Peruzzi et al. (22) presented the application of Multi-outcome Meshed Gaussian Processes on Projected Inputs (PIMGP). This method adapts Meshed Gaussian Processes, a method from the geostatistical literature which normally works with bidimensional inputs, for use in higher dimensional input spaces: after projecting the inputs onto a lower dimensional subspace (using, *e.g.* PCA), PIMGP use common GP kernels and lead to much faster performance relative to standard GPs or Bayesian Kernel Machine Regression (BKMR), especially with big data-frames (high N). The modeling framework was very flexible, allowing multiple outcome variables, missing covariates and covariates measured with error.

McGee et al. (1) presented the application of Bayesian Multiple Index Models (BMIM) (McGee et al., 2021). This approach combines the dimension reduction and interpretability of linear index models (such as WQSR and BGIR) and flexible exposure-response modelling of response surface methods (such as BKMR and PIMGP). With BMIM, the original exposures are reduced to a set of indices, as in BGIR. The approach simultaneously estimates the index component weights (with variable selection) and a potentially complex, high-dimensional exposure-response relationship between the indices and health outcome. Thus, it allows non-linear effects of the indices and interactions between indices. This presentation won one of two exposome data challenge prizes.

4.2. Approaches using Machine Learning to maximize prediction performance

Guimbaud (14) et al. linked the exposome with several health outcomes using several machine learning methods, namely multilayer

perceptron, random forest, XGboost, support-vector machines (SVM), and elastic net. They compared the prediction performance of the different techniques and used explainable AI by calculating SHAP (SHapley Additive exPlanations) values (Lundberg and Lee, 2017) to examine the impact of each exposure in the resulting models and their interactions with regards to the health outcomes. Zou (13) et al. conducted similar analyses, using in this case a deep neural network ensemble model, which was compared in terms of prediction accuracy to LASSO, SVM and random forest. Instead of SHAP values, they calculated a permutation-based feature importance test. Broséus (16) et al. studied the relationship between the exposome and child's BMI using a multi-step approach. First, they ranked predictors by feature importance using multivariate ordinal random forests. Based on this metric, they used an arbitrary threshold to select the most pertinent exposures and, among these, they performed an analysis of exposure associations using: 1) an ordinal logistic regression model to obtain effect estimates and direction of association and 2) an MCA to obtain a graphical view of the clustering of exposures, adapted to categorical exposures. Finally, Yun Chen et al. (24) proposed a measure of explained variation to assess the performance of models, which can be estimated accurately without estimating the individual regression coefficients. Looking at explained variation can provide interesting insights, such as confidence intervals, the explanatory power of different exposure families or of interactions terms.

4.3. Multi-stage approaches for combined effects

Multi-stage modelling approaches were applied to model patterns in the exposome data prior to examining associations with the outcome. For example, Pearce et al. (12) linked the exposome to birth weight by applying the framework defined as Exposure Continuum Mapping (ECM). An exposure continuum map is a spatially organized map of exposome features that places similar exposure profiles close to each other and different ones are further apart. It is built in two steps: first they build a low dimensional (2D) representation of the data using Kohonen self-organising maps (Kohonen, 1982) in order to identify exposure profiles. Then, using information from this organised map, they used a Generalized Additive Model (GAM) to build a 3D exposure-response function that allows examination of a total mixture effect.

Benavides et al. (8) linked the urban exposome with neurobehavior using a strategy that involved reducing the exposome via PCP (Gibson et al., 2021) and FA, in order to identify both consistent and unique pre- and postnatal exposure patterns, and then regress these lower dimensional patterns to the health outcome using generalized additive models (for the consistent pre- and postnatal patterns) and LASSO (for the unique patterns). Mistry (11) et al. linked the exposome to obesity. They used PCA and k-means clustering to identify exposure profiles in both the prenatal and postnatal periods, and then used logistic regression to assess the risk of obesity as a function of the transitions of individuals between prenatal and postnatal exposure clusters. Basu (23) et al. designed an iterative algorithm (COLRNS) that creates clusters of correlated exposures and then performs variable selection within the clusters to predict the health outcome while minimizing the error of the model. This method can also handle missing data.

4.4. Studies using omics data to improve inference on the link between exposome and health

Several studies used one or more omics datasets as intermediate layers and conducted some kind of mediation or meet-in-the-middle analyses. Wang (Ziyue) et al. (10) studied the link between the exposome and asthma using transcriptome as an intermediate layer. In particular, they used a combination of 1) differential gene expression and gene set enrichment for asthma, 2) exposure selection in a model for asthma via elastic net and calculation of exposure risk scores, and 3) high-dimensional mediation analysis. Shen et al. (18) also conducted

mediation analysis, in this case using several omics datasets (transcriptome, proteome, serum/urine metabolome) as potential mediators in the relationship between the exposome and several health outcomes (IQ, behavior, BMI). All models were fitted with linear mixed models and they used BH correction to correct for multiple comparisons. This presentation received the committee prize for integrating all the high dimensional omics and multiple outcomes while using informative visualisation of the results. Wang (Congrong) et al. (19) conducted a causal mediation analysis using multi-omics layers (transcriptome, proteome, serum/urine metabolome) as potential mediators of the relationship between the exposome and BMI. In this case, they used multi-omics factor analysis to reduce dimensionality of the omics layers and factor analysis to reduce the dimensionality of the exposome.

Yu et al. (21) presented Gatekeepers (Yu et al., 2021) a new theory to assess exposure-metabolites associations. They identified some Gatekeepers in the data using the Pearson correlation and then studied their associations with asthma. Gatekeepers are metabolites associated with both exposures and other metabolites. They are presented as the bridge between the exposome and the metabolome. Brunius et al. (17) also implemented a meet-in-the middle approach in which they used the proteome, serum metabolites, urine metabolites, gene expression and methylation as middle layers between the exposome and health outcomes. In their analytical pipeline, Omics Modules for Exposome Health Associations (OMEXA) (Shi et al., 2019), they used machine learning (MUVR Multivariate Methods with Unbiased Variable Selection, a predictive machine learning algorithm using an embedded recursive feature elimination mechanism within a repeated double cross validation procedure) to select the exposure variables related to the phenotypes available and partial correlation to further refine the selected list of exposures, adjusting for covariates. Then, they reused the same pipeline to select omic variables related to the selected exposures; and finally, they implemented a generalized linear model to link the selected omics with the phenotypes. They visualized the final results with triplots by projecting exposures, omics and phenotypes into two principal components.

Zhao (Stratakis) et al. (9) studied the link between organochlorines and BMI, using proteins, urine and serum metabolites as intermediate layers. They used the latent unknown clusters (LUCID) method (Peng et al., 2020), which found latent subgroups of subjects characterized by having at the same time distinguished BMI, distinguished omics profiles and distinguished exposure. The process includes variable selection with LASSO and results were visualized with a Sankey diagram.

Finally, Wu et al. (7) linked the exposome with metabolomics, without including the health phenotypes. They used a bipartite graph model to represent pairwise associations between exposures and metabolites. From this graph they extracted subgraphs of concentrated most significant negative or positive association blocks (that were assessed using the pairwise Pearson correlation coefficients).

5. Causal inference

Some researchers studied causal relationships between environmental exposure and health. Roscoe et al. (25) studied the relationship between air pollution and cognition, and used blood concentrations of metals (part of the exposome) as potential mediators. They used causal mediation analysis and quantile g-computation to assess mediated effects. Zugna et al. (20) also conducted mediation analyses, in this case they considered the exposome as a potential mediator in the association between socioeconomic position and birthweight. Thus, they studied a context with a high-dimensional mediator set, and their proposed analysis was based on interventional effects and penalized regression models (Loh et al., 2020).

Cáceres et al. (15) conducted an analysis trying to explain the differences in BMI between boys and girls (outcome variable). They inferred groups (clusters) of participants with specific exposomic profiles for which those differences were the highest. The underlying

method used was a recent implementation (<https://github.com/teff-pac/kage/teff>) of random causal forests (Wager and Athey, 2017). They also looked for omics markers (transcriptomic and methylomic) that were associated with differences between the exposome clusters. This presentation won the popular vote of the challenge.

6. Results on chemical pollutants and zBMI

In this section, we summarise the results obtained by different participating groups that addressed a similar research question, namely the relationship between chemical pollutants and zBMI. This is done for illustrative purposes and to provide some idea on how similar are the results when different methods are applied to the same dataset. The results of such comparison are of course dependent on the data and methods used and cannot be generalised, but they can give an idea of what could happen in a real research setting if multiple statistical analyses are applied. We also note that the dataset used was partially simulated, and that the analyses did not necessarily adjust for all necessary confounders. Therefore, subject-matter associations reported in this section should not be interpreted, and we just restrict the focus on the reproducibility of associations under the different analyses.

Although subject-matter interpretation of the results is not the focus of this section, it is noteworthy that most of the approaches that focused on child BMI confirmed previous HELIX publication results (Vrijheid et al., 2020), namely that childhood hexachlorobenzene (HCB) exposure is cross-sectionally associated with reduced childhood BMI z-score. Some studies also identified metals and PCBs (polychlorinated biphenyls, particularly PCB170) to be linked with BMI.

In a WQS regression of 38 prenatal and postnatal chemicals, Gennings et al. (4) reported a significant negative association between the mixture index and child BMI. Higher weights belonged to the postnatal exposures, with HCB being the chemical with the highest contribution. Conversely, Midya (2) et al. found in a penalized group mixture BWQSR that prenatal organochlorine compounds (OCs) and metals were positively associated with child BMI, whereas PCBs were negatively associated. Within each group, the chemicals with the highest weight were: HCB (for OCs), As, Cd and Co (for metals), and PCB170 (for PCBs).

Mork and Wilson (3) developed treed distributed lag mixture models with 56 prenatal and postnatal exposures and observed that the chemicals with the highest PIPs (near 1) were HCB, PCB170, DDE, and Mo. They also identified a strong interaction ($PIP = 1$) between prenatal Mo and postnatal HCB. McGee et al. (1) grouped 150 exposures into 29 indices corresponding to exposure families and time of exposure (prenatal or postnatal) in a BMIM. The groups with the highest PIPs (>0.5) were postnatal OCs, postnatal metals, prenatal water DBPs and the postnatal built environment. Among these, OCs were strongly negatively associated with child BMI z-score, and HCB was the chemical with the strongest effect in the index, followed by PCB170 and DDE. No interactions were observed except for prenatal water DBPs and postnatal OCs. Consistently, Broséus et al. (16) also identified postnatal HCB as the chemical exposure with the highest importance in multivariate ordinal random forests. After combining lifestyle and chemical exposures in an MCA, they found that Cu was associated with an increased risk of childhood overweight.

Using the LUCID method, Yinqi Zhao et al. (9) identified protein signatures (IL-1beta, IL-6, insulin) giving insight into underlying mechanistic pathways of childhood obesity (e.g. systemic inflammation, disturbed glucose metabolism). Finally, in multi-omics mediation analysis, Wang (Congrong) et al. (19) detected urine metabolites (e.g. phospholipids, TMAO, hippurate) that mediated the effect of maternal smoking and built environment on childhood BMI.

7. Discussion

This event brought together researchers from various disciplines to work on a common challenge: exposome data analysis. It established an

overview of state-of-the-art methods currently used in the field and paved the way to interesting discussions and exchange of ideas. Next, we discuss some of the advantages and disadvantages of the different methods proposed.

7.1. Dimensionality reduction

The first point of interest was the wide use of dimensionality reduction methods, especially feature selection, to deal with multivariate exposomic data (even without omics). Feature extraction methods were less used because they usually complicate the interpretation of the results if we are interested in the effect of a particular exposure on health. However, during the challenge, some interesting methods tried to analyse groups of correlated exposures as a way to reduce the dimensionality of the input while keeping the results interpretable.

7.2. Combined effects of exposures

In this section we discuss general pros and cons of the different methods presented. Several index methods were presented. These have the advantage of easy interpretation, as they provide a single parameter estimate for the mixture of exposures, along with the weights that easily illustrate the contribution of each exposure. The detection of a mixture effect is also expected to be more powerful when it is based on a single degree of freedom test. The weighted quantile sum regression family imposes directionality constraints, which could be seen as a limitation compared to other approaches such as Bayesian Group Index Regression. However, indices with both positive and negative weights are more difficult to interpret, and within the context of WQSR one can build one index for those exposures with positive contribution and another for those with negative contribution. Approaches such as LBWQSR may facilitate the interpretation of results by shrinking some of the less relevant associations towards zero. In principle, index models assume linear associations, but methods have been developed to estimate quadratic associations with the weighted index, where the significance of the quadratic term may be used as a test for the linearity assumption.

All index methods have the disadvantage that they do not consider interactions between exposures contributing to the same index. This can be solved by using response surface methods, at the cost of a potentially more difficult interpretation. Two methods were presented that searched for and detected two-way interactions in their analyses, and other methods such as PIMGP or BKMR can potentially capture higher order interactions and nonlinear effects. As order of interactions increases, flexibility increases, but again possibly at the price of interpretability. It is worth noting that, even if models can capture complex high-dimensional surfaces, interpretation of such models is usually done using plots that show the effects of just one or two variables at a time. Even though one could explore the effects of higher order interactions in methods such as PIMGP or BKMR, some studies suggest that structures based on four variables are at the limit of human ability to correctly process the information (Halford et al., 2005). Thus, when interest is in explaining the associations (and not merely predicting), higher order interactions can be of limited use.

The tension between interpretability and complexity when choosing between index models and response surface models can be eased by recently developed methods (multiple index models) that combine some of the advantages of both families of methods, and while defining easily interpretable indices, they can accommodate non-linear and non-additive relationships between exposure indices and the health outcome (McGee et al., 2021).

Several Bayesian techniques were presented in this section. The Bayesian paradigm is useful because it naturally penalises complex models and it offers flexibility to incorporate a process of variable selection. It is also useful to obtain the distributions of any quantity that can be derived from the model output. These techniques require some familiarity with Bayesian methods, for example to evaluate

convergence. However, currently available R packages greatly facilitate their implementation, even for those less versed in Bayesian inference.

7.3. Machine learning and prediction

Machine learning methods can potentially increase the predictability of the outcome by capturing more complex information from the data (e.g., complex interactions, non-linear relationships etc.). Models that combine multiple statistical techniques into an ensemble can even provide better results, as the different methods may be able to capture different patterns of the data. In the challenge, black box methods included ensemble methods (such as random forests, xgboost), neural networks and support-vector machines. The method most often used was by far random forests. One has to consider that the small sample size available in the challenge ($N = 1301$) limited the use and performance of machine learning methods.

Despite their potentially greater predictive power, machine learning techniques have been rarely used until recently in the context of environmental health studies, probably because of their lack of interpretability. One can improve on the interpretability of black box models in several ways, here we will discuss two of them. The feature importance metric was the most popular technique during the challenge and is also the most used in the exposome field. It can be computed in several ways depending on the model: 1) impurity based feature importance (Breiman, 2001) for a model based on decision trees 2) permutation based feature importance (Altmann et al., 2010) and shapley values based feature importance (Harris et al., 2021) can be both applied on any model (model agnostic). Another approach is to use partial dependence plots (Zhao and Hastie, 2021) that allow visualizing partial associations between variables. There are other approaches not implemented in the challenge that make use of machine learning while trying to obtain interpretable results. One example is the application of a combination of super-learner and g-estimation to assess the association between chemical pollutants and cognitive function (Oulhote et al., 2019).

7.4. Integration of omics data

Considering the analysis of omics data, all the studies presented performed some sort of dimensionality reduction before applying different statistical analysis. This is a limitation of most existing multi-omics data integration approaches since they have not been implemented to deal with large matrices. Therefore, development of new integrative methods/tools must consider efficient handling of large data sets (Subramanian et al., 2020). Most presentations studied the relationship between environmental exposures and health outcomes using omic data (single or multi-omics) as an intermediate layer in a mediation analysis fashion. Some other studies used Pearson correlation to study the relationships between omics and exposome data. Different combinations of statistical tools were proposed for omics data integration in analytical frameworks. These tools taken individually were not novel but their combination for integrating exposome, multi-omics and outcomes was strongly relevant and novel. We note that when one wants to include multi-omics layers in the analysis, the analyst needs to make several decisions in terms of pre-processing the data, reducing dimensionality, testing association between multiple sets of data using different techniques, and so on. This can lead to a large number of potential pipelines to analyse such data, each leading to potentially different results. This is a common problem in most kinds of data modelling, but it is aggravated in multi-omics analyses due to the availability of multiple sets of high-dimensional data.

Moreover, we note that other methods previously used for exposome and omics data may also be of interest when analysing such data, but they were not presented during this event. These methods include dimension reduction techniques related to PCA, such as partial least square (PLS) and its derivatives (sparse-PLS, group-PLS) (Chun and Keleş, 2010; Jain et al., 2018; Lenters et al., 2015), canonical-based

methods or network analysis (Bessonneau et al., 2021).

8. Causal models

Causal models have gained popularity in environmental epidemiology (Bind 2019), including for mixtures (Bellavia et al. 2019). Indeed, causal questions are what ultimately drive interventions and policy change. Causal mediation analysis with exposome data can help us prioritize environmental factors that have the greatest impacts on health. In the challenge, examples of causal models applied to exposome-health associations included mediation analysis using omics data, g-computation methods and the use of causal random forest.

With regards to mediation analysis with omics data, we note that our data was cross-sectional. In such a setting, results from mediation analyses with omics data should be interpreted with caution, since the omics markers might be a consequence of the health outcome or of the exposure (one or the other, not both). In particular serum metabolome data, which mainly included information on lipid metabolism, are closely related to phenotypic outcomes such as BMI. Therefore, the associations found should be expected to reflect more outcome classification (e.g. obesity subtypes) than an exposure effect.

The methods that used other causal inference methods focused on a clearly defined question that involved a single main variable of interest (sex or socioeconomic position) and its relationship with a health outcome, although they used the exposome as an intermediate layer. This is expected, as causal inference requires a clearly formulated causal quantity of interest that is easier to define for a single variable. Causal inference methods for a high-dimensional exposome is a field in which further developments are needed, possibly with a comparison of the results of those causal techniques with the results of agnostic analyses that perform variable selection, effectively treating all exposures in the same way.

9. Final remarks

We note that this article does not provide an exhaustive list of potential methods to study the exposome. In addition, the provided dataset was limited in power to study interactions between exposures. Still, this is a realistic setting for most epidemiological datasets with exposome data. Other approaches to be explored in the future include trying to integrate a priori knowledge in the analysis, for example from experimental toxicological data, in order to find chemical interactions, or to a priori group exposures with similar biological targets. These knowledge-driven approaches can also be used to reduce the dimensionality in the omics dataset. Other future challenges to be addressed in the exposome field include developing analysis strategies for longitudinal exposome datasets. The longitudinal component is actually key in the exposome definition, as the exposome tries to capture the totality of exposures across a lifespan. The present dataset only included two periods, pregnancy and childhood, and thus it only partially covered the lifetime exposome.

The strength of this event was the application of various methods on the same, well-characterized HELIX dataset. Indeed, it was possible for approaches focusing on the same outcome, child BMI, to find the same chemicals as main predictors (PCBs, metals). Although biological interpretation from these analyses should be avoided due to the partially simulated nature of the data and the heterogeneity in the methods to deal with the confounder structure (e.g., multi-centre structure), this exposome dataset could serve as a reference to test novel methods in the future. We also acknowledge that the way we imputed part of the data may have diluted some complex association patterns that may be present in the real dataset. Although we tested some models and obtained similar results in the real and imputed dataset, this may not be the case with the most flexible models, which can potentially capture complex association in the real dataset but not in the simulated dataset. Finally, the work herein has resulted in computational algorithms with

associated code made available to the community with an open-source licence, allowing for reproducible research and applications to other similar research questions based on exposome and even multi-omics data. This event was thought-provoking and highlighted the importance of networking between researchers in a multi-disciplinary environment. It fostered new collaborations at a time where interpersonal interaction was constrained by COVID pandemic, as well as giving visibility to researchers of various genders, backgrounds and career stages.

10. Consortia

The Exposome Data Challenge Participant Consortium is listed below in alphabetical order, and author affiliations are available in [Table S1](#).

Alfano Rossella, Basu Sanjib, Benavides Jaime, Broséus Lucile, Brunius Carl, Caceres Alejandro, Carli Matthew, Cazabet Rémy, Chattopadhyay Shounak, Chen Yun Hua, Chillrud Lawrence, Conti David, Gennings Chris, Gouripeddi Ramkiran, Iyer S Hari, Jedynek Paulina, Li Huichu, McGee Glen, Midya Vishal, Mistry Sejal, Moccia Chiara, Mork S Daniel, Pearce L. John, Peruzzi Michele, Pescador Jimenez Marcia, Reimann Brigitte, Roscoe J. Charlotte, Shen Xiaotao, Stratakis Nikos, Wang Ziyue, Wang Congrong, Wheeler David, Wilson Ander, Wu Qiong, Yu Miao, Zhao Yinqi, Zou Fei, Zugna Daniela.

CRedit authorship contribution statement

Conceptualization: L. Maitre, X.Basagaña, J.R. Gonzalez. Investigation (committee): X. Basagaña, C. Warembourg, L. Maitre, J.R. Gonzalez, J.B. Guimbaud, M.C. Hyam, P.M. Petrone. Data curation: X. Basagaña, C. Warembourg, L. Maitre, J.R. Gonzalez, M. Vrijheid. Writing - Original Draft: J.B. Guimbaud, X.Basagaña, L. Maitre. Formal analysis: all the event participants/consortia. Writing - Review & Editing: All authors. Funding acquisition: L. Maitre, M. Vrijheid.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

The data challenge data are fully available

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Appendix A. Supplementary material

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