



Research Article

Risk-Based Prioritization of PFAS Using Phenotypic and Transcriptomic Data from Human Induced Pluripotent Stem Cell-Derived Hepatocytes and Cardiomyocytes

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Abstract

Per- and polyfluoroalkyl substances (PFAS) are chemicals with important applications; they are persistent in the environment and may pose human health hazards. Regulatory agencies are considering restrictions and bans of PFAS; however, little data exists for informed decisions. Several prioritization strategies were proposed for evaluation of potential hazards of PFAS. Structure-based grouping could expedite the selection of PFAS for testing; still, the hypothesis that structure-effect relationships exist for PFAS requires confirmation. We tested 26 structurally diverse PFAS from 8 groups using human induced pluripotent stem cell-derived hepatocytes and cardiomyocytes, and tested concentration-response effects on cell function and gene expression. Few phenotypic effects were observed in hepatocytes, but negative chronotropy was observed in cardiomyocytes for 8 PFAS. Substance- and cell type-dependent transcriptomic changes were more prominent but lacked substantial group-specific effects. In hepatocytes, we found upregulation of stress-related and extracellular matrix organization pathways, and down-regulation of fat metabolism. In cardiomyocytes, contractility-related pathways were most affected. We derived phenotypic and transcriptomic points of departure and compared them to predicted PFAS exposures. Conservative estimates for bioactivity and exposure were used to derive a bioactivity-to-exposure ratio (BER) for each PFAS; 23 of 26 PFAS had BER > 1. Overall, these data suggest that structure-based PFAS grouping may not be sufficient to predict their biological effects. Testing of individual PFAS may be needed for scientifically-supported decision-making. Our proposed strategy of using two human cell types and considering phenotypic and transcriptomic effects, combined with dose-response analysis and calculation of BER, may be used for PFAS prioritization.

Plain language summary

Per- and polyfluoroalkyl substances (PFAS) are man-made chemicals used in many products. However, most of these substances have not been tested for safety, and concerns exist that they may be harmful to human health and/or the environment. This study aimed to use human cell-based models to investigate if some of the PFAS may exhibit hazardous properties and if similarities among substance groups are observed. Few effects were observed in liver cells, but a decrease in beating frequency was observed in heart cells for some PFAS. Gene expression changes were substance- and cell type-dependent. We did not find convincing structure-based similarities among PFAS; this suggests that testing of individual PFAS may be necessary in the future to inform health decisions. Overall, this study showed that a test strategy of using two human cell types, from liver and heart, may inform PFAS prioritization without a need for testing in animals.

1 Introduction

Per- and polyfluoroalkyl substances (PFAS) are man-made chemicals that contain carbon-fluorine bonds with distinct end functional groups (Carlson et al., 2022). The Organisation for Economic Co-operation and Development (OECD) maintains an inventory of

PFAS that contains over 4,700 substances (OECD, 2018), albeit estimates exceeding 10,000 PFAS have also been reported (Williams et al., 2017). PFAS are widely used in industrial applications and consumer products, and many of them are highly persistent, water soluble and mobile, leading to environmental dispersal and accumulation in the water, soil, wildlife, and ultimately humans.



While some PFAS, such as perfluorooctanoic acid (PFOA) and perfluorooctane sulfonic acid (PFOS), have been studied extensively and reports of their adverse health effects are abundant (Fenton et al., 2021; Meneguzzi et al., 2021), the vast majority of PFAS lack toxicity data. For example, a systematic evidence map of available epidemiological and animal bioassay data for a set of ~150 PFAS that were prioritized in 2019 by the US EPA for *in vitro* toxicity and toxicokinetic testing showed that only 45 compounds had some data in either animal or epidemiology studies (Carlson et al., 2022). However, a growing body of evidence suggests that other PFAS may also pose human health hazards (NTP, 2019a,b). Because of the growing concern about the persistence of PFAS and the potential that they may be hazardous to both human health and the environment, health and environmental protection authorities in Denmark, Germany, the Netherlands, Norway and Sweden proposed restrictions on PFAS as a class on 13 January 2023 (ECHA, 2023). Because of the considerable data gaps on virtually all PFAS, there is a significant need for and interest in accelerating the process of generating toxicology and exposure data that will inform decision-making.

One of the key needs in PFAS risk assessment is a scientifically-supported strategy for prioritization of PFAS for testing. In both the United States and Europe, regulatory bodies proposed a pragmatic approach of grouping PFAS into categories followed by selection of representative compounds for further study (ECHA, 2023; US EPA, 2021). A common approach is to group PFAS based on structural similarities (Buck et al., 2011; Sha et al., 2019; Wang et al., 2017; OECD, 2018). This approach has been applied to define groups of PFAS for risk assessment (Patlewicz et al., 2019; Buck et al., 2021). For example, the US EPA has established a workflow for PFAS evaluation that prioritized a subset of compounds to maximize information to support read-across within structure-based groupings (Patlewicz et al., 2019; Carlson et al., 2022).

The canonical read-across approach assumes that the chemical's toxicological properties can be inferred from those of a structurally similar chemical (Ball et al., 2016). Structure-property relationships are used to estimate the bioaccumulation potential of novel and emerging PFAS, as well as their protein binding and elimination rates (Cousins et al., 2020; Cheng and Ng, 2018; Gomis et al., 2018). Structure-property relationships have been reported for human hepatocyte lipid accumulation and gene expression in a study of 19 PFAS (Marques et al., 2022). Read-across approaches are already applied for PFAS management; for example, the State of Massachusetts proposed maximum contaminant level values for drinking water based on similarities in chemical structure and effects (Massachusetts Government, 2020). Still, questions remain whether grouping of PFAS based on structure alone is scientifically justifiable.

Additional data, including that from *in vitro* studies, may provide a biological dimension for informing class-based approaches for read-across and/or within-group prioritization for additional testing. Indeed, toxicity and toxicokinetic data are being generated using *in vitro* testing and computational methodologies to inform PFAS hazard characterization (US EPA, 2019;

Patlewicz et al., 2019, 2022; Kreutz et al., 2023; Dawson et al., 2023; Carstens et al., 2023). In addition, gene expression data offer a comprehensive understanding of chemical-induced bioactivity, delivering not only mechanistic insights but also dose-response information (Reardon et al., 2023; Johnson et al., 2022). Several studies have tested effects of PFAS on gene expression in human liver spheroids and demonstrated that both mechanistic insights and dose-response information can be used for ranking/prioritization and comparison to the effects in rodent liver (Rowan-Carroll et al., 2021; Reardon et al., 2021).

Based on the regulatory need to group PFAS and previous studies demonstrating the value of *in vitro* studies for dose-response analysis, we tested 26 PFAS from 8 structurally diverse groups in two human cell types from organs that are known to be targets for PFAS. We evaluated the hypothesis that structure-based grouping of PFAS can be substantiated using *in vitro* bioactivity data from both phenotypic and transcriptomic endpoints, and that these data may be also used for risk-based prioritization of PFAS. To address this hypothesis, we tested concentration-response effects on both cell function and gene expression in human induced pluripotent stem cell (iPSC)-derived hepatocytes and cardiomyocytes. The selection of these two cell types was informed by current knowledge and research gaps. Liver toxicity is one of the most studied effects of PFAS exposure (NASEM, 2022), while the potential cardiotoxic effects of PFAS exposure are currently understudied (NASEM, 2022), underlining the need for further investigation using *in vitro* models (Burnett et al., 2021a). Phenotypic and transcriptomic points of departure (POD) were derived to quantify the bioactivity of tested PFAS, and a transcriptomic pathway enrichment analysis was performed to determine potential similarities in the molecular effects of substances within and between chemical classes. Finally, the transcriptomic and phenotypic PODs were compared to exposure data to derive bioactivity-to-exposure ratios (BER).

2 Materials and methods

Chemicals and biologicals

Test PFAS ($n = 26$, Tab. 1) were obtained from the US National Toxicology Program as 20 mM stocks in 100% tissue culture grade dimethyl sulfoxide (DMSO). The PFAS were selected based on preliminary bioactivity data, from which high and low rank chemicals were then selected in a smaller subset of PFAS with diverse structures for additional *in vitro* screenings. The PFAS were assigned into sub-categories using the head-group/side-chains to determine their classification as previously suggested (OECD, 2018; Buck et al., 2011, 2021). Tested PFAS, abbreviations, and their chemical structure-based category information are listed in Table 1. Tissue culture grade DMSO (CAS# 67-68-5, sc-358801) was used as the vehicle for all studies (Santa Cruz Biotechnology, Dallas, TX, USA).

Cell-specific positive controls were used for both the iPSC-derived hepatocytes (iPSC-Hep) and cardiomyocytes (iPSC-CM). The positive controls for the iPSC-Hep were rotenone (CAS# 83-79-4, Cat# 45656-250MG), amiodarone (CAS# 19774-82-4,



Tab. 1: PFAS tested in this study

CASRN	Chemical name	Chemical abbreviation	Category	Category abbreviation
2043-47-2	4:2 Fluorotelomer alcohol	4:2 FTOH	n:2 Fluorotelomer alcohols	n:2 FTOH
678-39-7	8:2 Fluorotelomer alcohol	8:2 FTOH		
375-73-5	Perfluorobutanesulfonic acid	PFBS	Perfluoroalkane sulfonic acids/ perfluoroalkane sulfonates	PFSA
2806-15-7	Sodium perfluorodecanesulfonate	PFDS-Na		
39108-34-4	8:2 Fluorotelomer sulfonic acid	8:2 FTS	n:2 Fluorotelomer sulfonic acids	n:2 FTSA
27619-97-2	6:2 Fluorotelomer sulfonic acid	6:2 FTS		
355-80-6	1H,1H,5H-Perfluoropentanol	PFPOH	Alcohols	Alcohols
335-99-9	Dodecafluoroheptanol	DDFHp		
423-54-1	Perfluorooctanamide	PFOAM	Perfluoroalkane amide/amines	PFAN
13485-61-5	Nonafluoropentanamide	PFNAM		
41997-13-1	Perfluorohexanesulfonamide	PFHxSA		
307-31-3	Perfluorooctanamide	PFOAMD		
356-02-5	3:3 Fluorotelomer carboxylic acid	3:3 FTCA	Fluorotelomer carboxylic acids	FTCA
335-67-1	Perfluorooctanoic acid	PFOA	Perfluoroalkyl carboxylic acids/ perfluoroalkyl carboxylates	PFCA
376-06-7	Perfluorotetradecanoic acid	PFTeDA		
72629-94-8	Perfluorotridecanoic acid	PFTIDA		
3825-26-1	Ammonium perfluorooctanoate	NH4PFOA		
335-76-2	Perfluorodecanoic acid	PFDA		
2058-94-8	Perfluoroundecanoic acid	PFUnDA		
375-95-1	Perfluorononanoic acid	PFNA		
1763-28-6	3,3-Bis(trifluoromethyl)-2-propenoic acid	TFPrOA		
375-85-9	Perfluoroheptanoic acid	PFHpA		
55621-21-1	Perfluoro-3,6-dioxaoctane-1,8-dioic acid	PFHx2Et2OA	Perfluoroalkyl ether carboxylic acids	PFECA
377-73-1	Perfluoro-3-methoxypropanoic acid	PFMPA		
330562-41-9	Perfluoro-3,6,9-trioxatridecanoic acid	PFPE-6		
863090-89-5	Perfluoro(4-methoxybutanoic) acid	PFMOBA		

Cat# A8423-1G), acetaminophen (CAS# 103-90-2, Cat# A5000-100G), and CCCP (carbonyl cyanide 3-chlorophenylhydrazide, CAS# 555-60-2, Cat# C2759-100MG); all were purchased from Sigma-Aldrich (St. Louis, MO, USA). Isoproterenol, sotalol, and propranolol (Molecular Devices, San Jose, CA, USA) were used as the positive controls for the iPSC-CM. Tetra-octyl ammonium bromide (TAB, CAS#14866-33-2, Cat# D2438) was obtained from Sigma-Aldrich (St. Louis, MO, USA) and used as a positive control for both cell types.

Human iPSC-Hep and iPSC-CM cells were obtained from FUJIFILM Cellular Dynamics (Cat# C1023 and Cat# C1006, respectively; Madison, WI, USA), along with their corresponding plating and maintenance media. Additional reagents includ-

ing RPMI medium (Cat# 11875-093), dexamethasone (Cat# A13449), B27 supplement (Cat# 17504-044), gentamicin (Cat# 15750-060), and penicillin-streptomycin (Cat# 15140-122) were obtained from Gibco (Waltham, MA, USA). Oncostatin-M was from R&D Systems (Cat# 295-OM-010, Minneapolis, MN, USA).

For cell culture, we used tissue culture grade 384-well plates (for iPSC-CM, Cat# 3764, Corning Life Sciences, Kennebunk, ME, USA) and collagen I pre-coated 384-well plates (for iPSC-Hep, Cat# 354664, Corning Life Sciences). Gelatin (Cat# G1890-500G), Trypan Blue 0.4% solution (Cat# T8154-100ML), and D-PBS (Cat# D8537) were obtained from Sigma-Aldrich.



Cell culture

Cells were plated and maintained according to instructions provided by FUJIFILM Cellular Dynamics and as described previously (Sirenko et al., 2014; Grimm et al., 2015). Briefly, iPSC-Hep were thawed for 3 min in a water bath at 37°C. The thawed cells were then transferred to a 15 mL tube; 9 mL of plating medium was then added. The plating medium contained RPMI medium with 2% (v/v) iCell Hepatocyte Medium Supplement, 0.1 μM dexamethasone, 2% (v/v) B27 supplement, 25 μg/mL gentamicin, and 20 ng/mL Oncostatin-M. The cells were counted using an automated cell counter and subsequently diluted to reach the final cell density of 7.2×10^5 cells/mL. The cell suspension was then transferred to collagen I pre-coated 384-well plates, yielding a final seeding density of 18,000 cells/well. Cells were plated in the inside wells (a total of 308 wells), and the outer wells were filled with PBS to help insulate the inner wells and avoid evaporation. Plates were kept at room temperature for 30 min before they were placed in the incubator at 37°C and 5% CO₂. Four hours post-plating, the plating medium was exchanged with 25 μL of fresh plating medium. Thereafter, the plating medium was exchanged daily until day 5 post-plating. On day 5, the plating medium was exchanged with maintenance medium made from RPMI medium containing 2% (v/v) iCell Hepatocyte Medium Supplement, 0.1 μM dexamethasone, 2% (v/v) B27 supplement, and 25 μg/mL gentamicin; this medium was used for the remainder of the experiment and replaced every other day until the chemical exposure on day 7.

For experiments in iPSC-CM, 384-well plates were prepared by adding 25 μL of 0.1% (w/v) gelatin solution per well and incubated for 2 h at 37°C and 5% CO₂. Vials containing iPSC-CM were thawed for 3 min in a water bath at 37°C. The contents of a single vial were then added to a 15 mL conical tube and 9 mL of plating medium containing 1:500 (v/v) penicillin/streptomycin solution was added. An aliquot of the cell suspension was used to count the cells, further dilutions were done to yield the target cell seeding density of 2×10^5 viable cells/mL. Immediately before plating, the gelatin solution was fully aspirated from the plates and 25 μL of cell suspension was added to each of the 308 outer wells, resulting in a cell density of 5,000 viable cells/well. The outer wells were filled with PBS to help insulate the wells and avoid evaporation. Plates were kept at room temperature in the hood for 30 min and then placed in the incubator at 37°C and 5% CO₂. Forty-eight hours after cell seeding, the plating medium was exchanged by removing 17.5 μL/well and replacing it with 32.5 μL of maintenance medium containing 1:500 penicillin/streptomycin. Maintenance medium was subsequently changed every other day for the remaining 12 days in culture until the chemical exposure and assays. On the evening before chemical addition, the medium was removed and replaced by 25 μL of fresh maintenance medium.

To evaluate the functional, cytotoxic, and genomic endpoints in iPSC-Hep, they were exposed to the PFAS for 48 h. The iPSC-CM were exposed to the PFAS for 90 min for functional and cytotoxic endpoints and for 24 h for the gene expression studies. For the functional and cytotoxic phenotypes, various stains and images were collected at the respective timepoints as detailed below, and for the genomic endpoints cell lysates were collected at the respective timepoints for transcriptomic analyses.

Plate design

A 200× chemical master plate was first prepared in 100% cell culture grade DMSO, the chemicals were then transferred to the 5× working plate (2.5% DMSO and 97.5% cell-specific culture medium). For the final assays (1×), 12.5 μL of the solutions in the working plate were added to the assay plate that contained cells in 50 μL of cell-specific medium, yielding a final concentration of 0.5% DMSO. Each assay plate contained 308 wells with cells and (i) each chemical (n = 26) and concentration (PFAS were tested in 4 final concentrations of 0.1, 1, 10, and 100 μM), (ii) vehicle (DMSO at final concentration of 0.5%) wells (n = 14), (iii) media-only wells (n = 6), and (iv) wells with cell type-specific positive controls (n = 20). The entire chemical layout was repeated twice per 384-well plate. The top tested concentration (100 μM) was not used for the genomic assays.

Phenotype assays

On day 7 post-plating, iPSC-Hep were exposed to test chemicals for 48 h. Chemical transfer from the master plate (200×) to the working plate (5×) and then to the assay plate (1×) was done using automated liquid handling of the FLIPR Tetra (Molecular Devices, San Jose, CA). After a 48-h exposure at 37°C and 5% CO₂, the medium was replaced with medium containing fluorescent dyes – MitoTracker, Hoechst 33342, and Calcein AM – to assess mitochondrial, nuclear, and cytoplasmic endpoints. Following a 15-min incubation at 37°C, the medium with dyes was replaced with fresh maintenance medium. The plates were then transferred to the ImageXpress Micro Confocal High-Content imaging system (Molecular Devices) for image analysis. Data for functional and cytotoxic endpoints was extracted using MetaXpress Software Multiwavelength Cell Scoring Module (Molecular Devices).

On day 14 post-plating, for the iPSC-CM, Ca²⁺ flux was measured before and after chemical exposure using the EarlyTox Cardiotoxicity Kit (Molecular Devices) as detailed in (Grimm et al., 2015). First, the Ca²⁺ dye reagent (25 μL) was added to each well and incubated at 37°C for 2 h. A baseline read was recorded using the FLIPR Tetra. Chemicals were then immediately added from the working plate (5×), transferring 12.5 μL to the assay plate (1×) (total volume of 62.5 μL) using the automated liquid handling of the FLIPR Tetra from the 5× working plate, the plates were then incubated for 90 min at 37°C. The Ca²⁺ flux was measured again, and the medium with chemicals was then replaced with the medium containing fluorescent dyes, specifically MitoTracker (2 μg/mL) and Hoechst 33342 (200 nM). After a 15-min incubation with the fluorescent probes, the medium was replaced with fresh maintenance medium and the plates were transferred to the ImageXpress Micro Confocal High-Content imaging system (Molecular Devices) for image acquisition. The imaging data were subsequently analyzed using the MultiWavelength Cell Scoring Module available on the MetaXpress software, and the data from relevant endpoints were extracted using both MetaXpress Software on the ImageXpress Micro Confocal High-Content Imaging System for cytotoxic endpoints and Screenworks 4.0 software on the FLIPR Tetra for the functional endpoints (Molecular Devices). The raw data for



Tab. 2: Cell types used and *in vitro* readouts collected in this study

Cell type	Phenotype	Exposure duration	Benchmark response for POD derivation
Human induced pluripotent stem cell-derived hepatocytes (iPSC-Hep)	Total cell number	48 h	1 SD
	Mitochondrial integrity		
	Mitochondrial intensity		
	Cytoplasmic integrity		
	Cell mean area		
	Gene expression	See methods	
Human induced pluripotent stem cell-derived cardiomyocytes (iPSC-CM)	Total cell number	90 min	10% change
	Positive chronotropy		5% change
	Negative chronotropy		5% change
	Asystole		95% change
	QT prolongation		5% change
	Gene expression	24 h	See methods

both cell types and all phenotypes are available as Supplementary Files 1¹ and 2².

Derivation of the phenotypic points of departure (pPODs)

For pPOD derivation, first, normalization of phenotypic readouts to their respective vehicle (0.5% DMSO) controls was conducted. The vehicle-treated (0.5% DMSO) samples were then screened to identify any outliers (measurements falling outside of the interquartile range), and outliers were removed if present. Vehicle control-scaled data for each test substance and phenotype were then fitted to a curve with a nonlinear logistic (Hill) function to determine POD values (Sirenko et al., 2017), defined as the concentrations at which the fitted curve exceeds certain thresholds above or below the mean of vehicle-treated controls (benchmark responses for each phenotype are detailed in Tab. 2). For iPSC-Hep, the choice of one standard deviation “benchmark response” was based on US EPA guidance (Wignall et al., 2014) and empirical observations that generate consistently high classification accuracy (Sirenko et al., 2017). For iPSC-CM, the POD values were derived based on the criteria consistent with human clinical cardiotoxicity phenotypes as previously described (Blanchette et al., 2020).

TempO-Seq library preparation and sequencing

Following chemical exposure, the wells were completely aspirated, 10 µL/well of 2× lysis buffer (BioSpyder Technologies, Carlsbad, CA) was added to each well, and plates were placed on an orbital plate shaker at room temperature for 10 min to facilitate cell lysis. Then, adhesive seals were placed on the plates and the lysates were frozen in the 384-well plates and stored at -80°C until further processing. The Templated Oligonucleotide Sequencing Assay (Tem-

pO-Seq™, BioSpyder Technologies) was used as the RNA sequencing technology, and gene expression was evaluated using the human TempO-Seq Whole Transcriptome panel (BioSpyder Technologies) consisting of 22,537 protein-coding probes. Detailed protocols for TempO-seq are provided by the manufacturer and have been previously reported (Grimm et al., 2016; House et al., 2017). Briefly, an aliquot (2 µL) of the cell lysate was transferred to a 96-well plate, in which the samples first underwent annealing to match detector oligos, followed by nuclease digestion of excess oligos, and detector oligo ligation, which were subsequently amplified with the tagged primers according to manufacturer instructions (BioSpyder Technologies). The amplified samples were then pooled and purified into a sequencing library using a PCR clean-up kit (Clontech, Mountain View, CA). The libraries were sequenced (single-end, 50 bp in length) using HiSeq 2500 v.2 (Illumina, San Diego, CA). The reads were then aligned to target probe sequences to generate a gene count matrix using the analysis pipeline detailed elsewhere (House et al., 2017) including the STAR aligner (Dobin et al., 2013) with the maximum number of allowed mismatches of 3, insertion/deletion open, and extension penalty of 0.

Gene expression data analysis

Probe read count data for transcripts with more than one probe in the TempO-Seq assay was first summed to gene level. Quality control steps with the following criteria were performed: 1) genes with fewer than 5 mean counts across the entire sample space (n = 330 for iPSC-Hep and n = 220 for iPSC-CM) were removed, 2) samples with mean expression across all genes lower than 25 were removed, 3) outliers in vehicle control samples were identified and removed using mean correlation at < 0.85 as the criterion to ensure the stability of further transcriptomic dose-response and differential

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² doi:10.14573/altex.2311031s2



expression analyses, 4) in addition, principal component analysis (PCA) was conducted on vehicle control samples using raw counts to identify any remaining outliers. After these quality control steps, 330 samples and 10,205 genes were retained for further analyses in iPSC-Hep, and 202 samples and 15,112 genes in iPSC-CM.

Bioconductor's DESeq2 package (Love et al., 2014) was used to perform differential gene expression analysis and to rank genes based on the log₂-fold-change values by contrasting the samples from the highest tested concentration (10 mM) of each PFAS substance and the vehicle controls. The ranked genes were then analyzed using Gene Set Enrichment Analysis (GSEA) approach (Subramanian et al., 2005) to derive enriched Reactome pathways (Jassal et al., 2020) as implemented in the gsePathway functions of R package ReactomePA version 1.16.2 (Yu and He, 2016). To enhance the interpretability, an additional data integration step was implemented to group significantly enriched gene sets from lower-level pathways into common higher-level nodes based on the Reactome pathway hierarchy (Jassal et al., 2020).

Analysis of variance (ANOVA) for categorical effects on individual genes was performed using the aov command in R. For those genes that were significant (FDR q-value < 0.05), a two-tailed post-hoc Tukey test was conducted using the TukeyHSD command in R as a guide to describe the categorical gene expression effects between individual PFAS subgroups.

Supervised category analysis using prediction analysis of microarrays

A machine-learning statistical model was trained to predict the structure-based PFAS categories using the Prediction Analysis of Microarrays (PAM) approach. The approach performs class prediction via nearest shrunken centroid for a high-dimensional vector and works for any quantitative set of features. The analysis utilized gene expression log₂-fold-change profiles as predictors, along with bioassay measurements (phenotypic PODs) in both cell types, performed using pamr version 1.56.1 (Tibshirani and Efron, 2002). PFAS categorical predictions were made using expression data alone and phenotypic PODs alone in each cell type, as well as a combination of expression data and phenotypic PODs in both cell types. For this analysis, the FTCA "group" with only one PFAS member was removed, as it is not suitable for cross-validation. The cross-validated matching accuracy, derived from the associated confusion matrix, was computed to show the proportion of matches between predicted category assignments using training data only, and true assignments. The proportion of matches expected under chance was determined by 1 million random permutations of group assignments, with a mean of 0.21 and 95th percentile of 0.32. Thus, a cross-validated matching prediction proportion in excess of 0.32 would be considered significant at the 0.05 level.

Transcriptomic benchmark dose modelling and derivation of transcriptomic points of departure (tPODs)

The Bioconductor's DESeq2 package (Love et al., 2014) was utilized to assemble the complete normalized count matrix as detailed

above. The normalized count data (counts +1 to zero protect the data for further analyses on the logarithmic scale) were processed using the BMDEExpress (v.2.3) software (Phillips et al., 2019) for transcriptomic benchmark dose modelling as detailed in (Tsai et al., 2023). Data were first pre-filtered with Williams' trend test (p-value ≤ 0.05 within each transcript and substance) and an absolute fold change ≥ 1.5 (compared with vehicle controls); genes that did not pass these criteria at any dose were removed from further analysis. Next, data were analyzed using Hill (version 2.18), power (version 2.19), linear (version 2.21), polynomial 2 (version 2.21), and exponential (2, 3, 4, and 5, version 1.11) models. A benchmark response of one standard deviation was used at the individual gene level to derive a benchmark dose (BMD). The best-fit model for each transcript was selected based on the following parameters: (1) maximum iterations of 250; (2) confidence level of 0.95; (3) constant variance; Hill models with a k parameter < 1/3 of the lowest positive dose were flagged and then the next best model with a p-value > 0.05 was used; and (5) a nested chi-squared cutoff value of 0.05 to select the best polynomial models followed by minimum Akaike Information Criterion value and a goodness-of-fit p-value > 0.05. The BMD output data files from BMDEExpress are available as Supplementary Files 3 and 4.

Transcripts that had BMD upper bound (BMDU) to BMD lower bound (BMDL) ratio > 40 or a model fit p-value < 0.1 were excluded from further analyses. Vehicle control samples were assigned a "dose" one log₁₀ unit below the lowest dose tested to allow plotting on the logarithmic scale, as well as allowing capping the BMD values at the lowest dose tested. In addition, BMD values were also capped at the highest dose tested.

For derivation of tPOD for each tested substance, the lowest gene set median gene BMD was used based on features that passed the criteria described above in accordance with previously published guidance (NTP, 2018). Gene sets were represented by both enriched pathways (Reactome and KEGG annotations) and Gene Ontology (GO) biological process and were identified using xgr package version 1.1.8 in R (Fang et al., 2016). The gene set background list included all interrogated genes retained after low count removal. Subsequently, the lowest median BMD value from any gene set was selected as the tPOD for each tested PFAS and cell type.

Selection of CRGs that were common across multiple PFAS

To assess whether different PFAS had similar concentration-response effects at the gene level, we employed permutation-based significance testing. Specifically, for each cell type separately, the binary recorded matrix was established, with individual CRGs as rows and PFAS as columns, where 0 indicated no significant concentration-response for each gene-PFAS pair, and 1 indicated that a PFAS elicited concentration-response of a transcript. Then, the data matrix was permuted column-wise, shuffling the elements of each column randomly. For each permutation iteration, the sum of each row (instances of a gene declared as CRG) across all PFAS was calculated. This process was repeated 10,000 times to construct a null distribution for each gene. The row sums from each

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⁴ doi:10.14573/altex.2311031s4

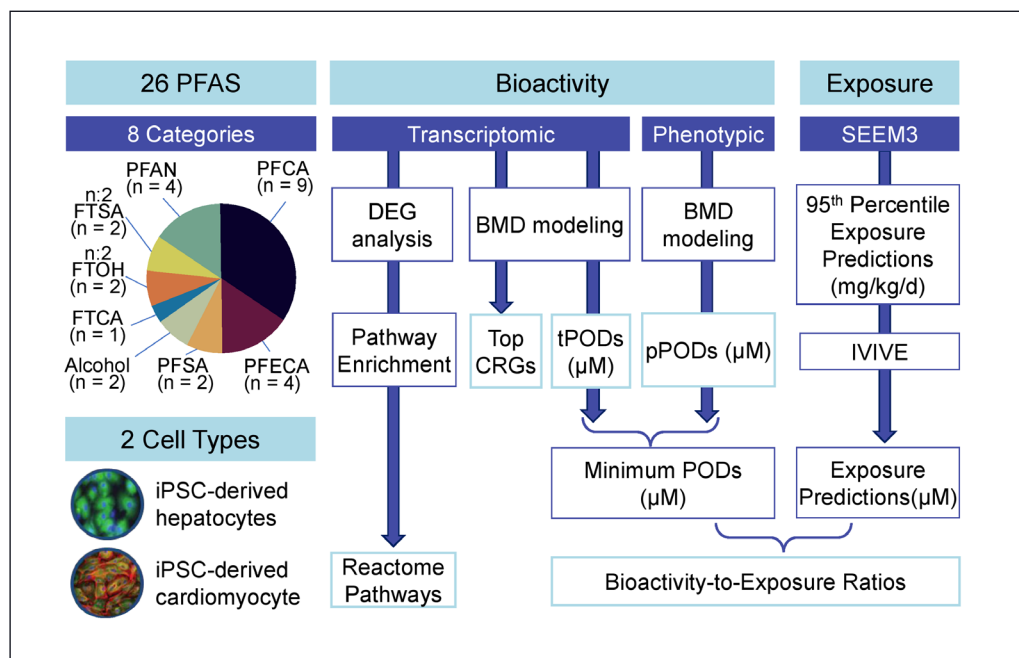


Fig. 1: Overview of the study design and data analyses of the *in vitro* effects of 26 PFAS in iPSC-Hep and iPSC-CM

See abbreviations for PFAS classes in Tab. 1. Other abbreviations: DEG, differentially expressed genes; BMD, benchmark dose; CRG, concentration-response gene; POD, point of departure (t for transcriptomic and p for phenotypic data); IVIVE, *in vitro*-to-*in vivo* extrapolation

permutation were stored for each iteration. The empirical p-value for each gene was then computed by comparing the observed sum for each gene (data from the actual experiment) to the null distribution. Genes with empirical p-values < 0.05 were considered statistically significant.

Derivation of the bioactivity-to-exposure ratios (BER)

To compare the bioactivity derived from *in vitro* assays and exposure levels, *in vitro*-to-*in vivo* extrapolation was performed using the equation detailed below (Eq. 1) to convert exposure dose level (mg/kg/day) to μM units. The equation was adopted from (Wetmore et al., 2012), where the original format was to convert an *in vitro* bioactivity value (μM) to an oral equivalent dose (mg/kg/day).

$$\text{Exposure estimates } (\mu\text{M}) = \frac{95^{\text{th}} \text{ percentile exposure prediction (mg/kg/day)} \times C_{\text{ss}} (\mu\text{M})}{1 \text{ (mg/kg/day)}} \quad (\text{Eq. 1})$$

This conversion used either human steady-state plasma concentrations (C_{ss}) from the htkk R package (Pearce et al., 2017) or the predicted fraction unbound data from recent publications (Kreutz et al., 2023; Smeltz et al., 2023). See C_{ss} and exposure values for 18 PFAS that had available data from either source in Table S1⁵. To ensure consistency and conservative assumptions when integrating C_{ss} data from various sources, intrinsic clearance was set to zero during the derivation of C_{ss} values. The BERs (Paul Friedman et al., 2020) were computed by dividing POD values from the *in vitro* experiments (this study, μM) by the 95th percentile exposure predictions (μM , as calculated using Eq. 1) for the general US population, as derived from the ExpoCast Systematic Empirical

Evaluation of Models version 3 (SEEM3) framework (Ring et al., 2019). The equation for BER calculation is given below (Eq. 2)

$$\text{Bioactivity-to-Exposure Ratio (BER)} = \frac{\text{Points of departure } (\mu\text{M})}{\text{Exposure estimates } (\mu\text{M})} \quad (\text{Eq. 2})$$

3 Results

This study tested the hypothesis that structure-based grouping of PFAS can be substantiated using *in vitro* bioactivity data from both phenotypic and transcriptomic endpoints and that these data may be used for risk-based prioritization of PFAS. The study's overall design and data analysis workflow are depicted in Figure 1. Specifically, we evaluated 26 PFAS (Tab. 1) that belong to 8 sub-groups based on established nomenclature (OECD, 2018; Buck et al., 2011, 2021). Bioactivity of PFAS compounds was tested using iPSC-hepatocytes (iPSC-Hep) and iPSC-cardiomyocytes (iPSC-CM), cells selected to be representative of organs of concern for potential adverse effects of PFAS (NASEM, 2022). Both phenotypic and transcriptomic data were analyzed to identify points of departure (PODs) and potential underlying mechanisms of PFAS effects. For risk characterization, we calculated bioactivity-to-exposure ratios (BERs) using conservative exposure estimates and the lowest POD.

3.1 Phenotypic effects of PFAS in iPSC-Hep and iPSC-CM

PFAS compounds (Tab. 1) were tested in iPSC-Hep and iPSC-CM in a range of concentrations (0.1, 1, 10, and 100 μM), and both functional and cytotoxicity phenotypes were evaluated (Tab. 2).

⁵ doi:10.14573/altex.2311031s8

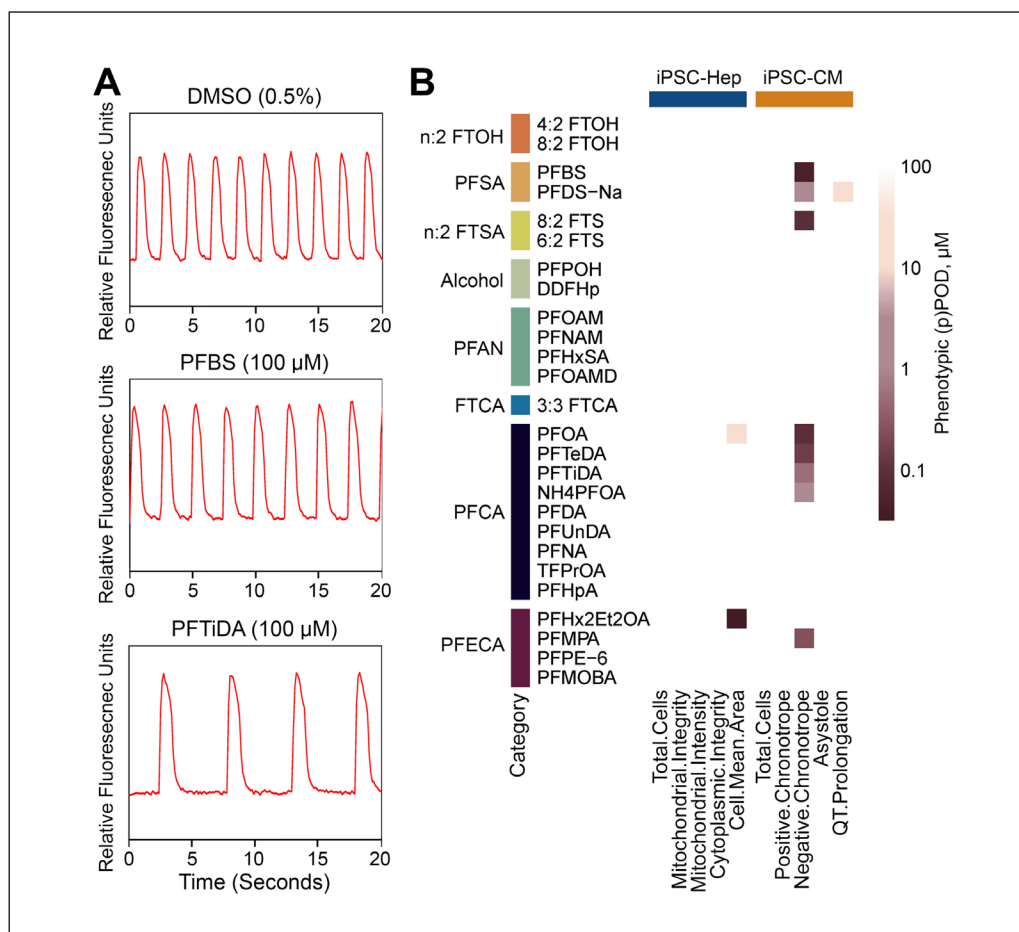


Fig. 2: Phenotypic effects of PFAS on iPSC-Hep and iPSC-CM

(A) Representative Ca^{2+} flux traces (90 min after treatment) for vehicle (0.5% DMSO), PFBS (100 μM), or PFTiDA (100 μM) in iPSC-CM. (B) Heatmap illustrating phenotypic points of departure (pPOD) for the phenotypes (columns) evaluated in iPSC-Hep (on the left) and iPSC-CM (on the right) for each tested PFAS (rows). A darker color indicates a lower POD value. PFAS were grouped by chemical structure categories. See Tab. 1 for class and individual chemical abbreviations.

Fluorescent imaging of the cytoplasm, nuclei, and mitochondria were used as relevant phenotypes in iPSC-Hep (Grimm et al., 2015; Sirenko et al., 2014). In iPSC-CM, Ca^{2+} flux was used as the primary indicator of the ion channel activity and beat frequency (Sirenko et al., 2013, 2017). Figure 2A shows representative Ca^{2+} flux traces for iPSC-CM that were treated with vehicle (spontaneous frequency of ~ 30 beats per minute) or two representative PFAS that elicited negative chronotropy effects at the highest concentration tested (100 μM) – perfluorobutanesulfonic acid (PFBS, 24 beats per minute) and perfluorotridecanoic acid (PFTiDA, 12 beats per minute). Dose-response modeling was performed on the data from each phenotype, and PODs were derived using phenotype-specific benchmark responses as indicated in Table 2.

Figure 2B depicts a heatmap of phenotypic PODs (pPOD) for each individual PFAS in both iPSC-Hep and iPSC-CM. In this supervised analysis, PFAS were arranged by their respective chemical structure group (Tab. 2), by cell type, and phenotype. In iPSC-Hep, only two PFAS elicited concentration-response phenotypic effects – PFOA and PFHx2Et2OA; both compounds increased mean cell area, which is indicative of cell enlargement most likely due to fatty acid accumulation. In iPSC-CM, 8 PFAS elicited effects that reached a benchmark response (Tab. 2) for two phenotypes – decreased peak frequency (PFBS, PFDS-Na, 8:2 FTS,

PFOA, PFTeDA, PFTiDA, NH4PFOA, and PFMPA), and QT prolongation (PFDS-Na). No patterns in phenotypic effects of tested PFAS in either cell type were evident with respect to PFAS chemical structure-based classes except for negative chronotropy, whereby there were relatively higher numbers of substances in PFCA and PFSA groups with effects; however, one-way ANOVA showed no significant difference among classes ($p = 0.22$). In addition, no significant correlation was found between pPOD and either chain length (Spearman $\rho = -0.15$, $p = 0.50$ in iPSC-CM; $\rho = -0.02$, $p = 0.94$ in iPSC-Hep) or molecular weight ($\rho = -0.29$, $p = 0.15$ in iPSC-CM; $\rho = 0.06$, $p = 0.76$ in iPSC-Hep) for the tested PFAS.

3.2 Transcriptomic effects of PFAS in iPSC-Hep and iPSC-CM

While few concentration-response phenotypic effects were observed at the whole cell-level, gene expression was evaluated to determine whether PFAS had molecular-level effects in both iPSC-Hep and iPSC-CM. These analyses (Fig. 1) included (i) determination of the effects at gene and pathway level at the highest concentration tested for gene expression analysis (10 μM), (ii) analysis of the concentration-response at the gene level (vehicle, 0.1, 1 and 10 μM), and (iii) derivation of the transcriptomic PODs at the pathway/gene set level (tPODs).

First, we used raw gene expression data for each PFAS (10 μ M) to conduct a principal component analysis (PCA). When all samples (PFAS and vehicle, in both cell types) were included in the PCA analysis (Fig. S1⁵), the greatest separation was observed between cell types, as expected. While vehicle-treated samples were most distinguishable from PFAS-treated ones in iPSC-Hep, some compounds exhibited effects in both cell types. Therefore, differential gene expression was analyzed separately by cell type and compared to vehicle-treated samples. Two PCAs are shown in Figure 3 to visualize such analyses for each cell type; individual PFAS are colored according to their chemical structure-based class (Tab. 1). This analysis showed that differential gene expression effects of PFAS exhibited little grouping by chemical structure-based class in both cell types. To further investigate the grouping effects from gene expression data, ANOVA analyses were performed on log₂-fold-change values for individual genes derived from comparing the highest concentration and control samples. Only a very few genes (9 out of 10,205) in iPSC-Hep were found to exhibit significant differences across the groups, whereas no gene showed significance in iPSC-CM. The results indicate minimal effects on expression corresponding to the structure-based categories in both iPSC-Hep and iPSC-CM. The list of genes and the between-group comparisons that passed the post-hoc Tukey tests are presented in Supplementary File 5⁶. Furthermore, to directly assess the potential correspondence of multiple features in gene expression and phenotypic bioactivity data to the structure-based groups, the PAM approach (see Section 2) was used to fit a machine-learning model for PFAS categorical predictions (Fig. S2, S3⁵). However, the prediction accuracy was not significant across all data types tested: transcriptomic data, phenotypic POD data, and a combination of both.

Second, as a surrogate measure of the potency of the tested PFAS' effect on transcription, we evaluated concentration-response for each compound by determining the number of concentration-responsive genes (CRGs) in each cell type (Fig. 4). Higher counts of CRGs may be interpreted as greater bioactivity. In iPSC-Hep, the number of CRGs affected by PFAS ranged from 116 (representing 1.1% of the total number of expressed genes in this cell type) to 542 (5.3%). In iPSC-CM, the numbers of PFAS-affected CRGs varied more widely, from 83 (0.5% of the total number of expressed genes in this cell type) to 2,027 (13.4%). The bar plots in Figure 4 show that the number of CRGs for individual PFAS were substance- and cell type-specific and not dependent on their chemical structure-based class ($p = 0.26$ in iPSC-Hep; $p = 0.75$ in iPSC-CM using one-way ANOVA). No significant correlation was found between the number of CRGs and either chain length ($\rho = -0.08$ in iPSC-CM; $\rho = 0.14$ in iPSC-Hep) or molecular weight ($\rho = -0.17$ in iPSC-CM; $\rho = 0.17$ in iPSC-Hep) for tested PFAS.

Third, we examined whether there were similarities in CRGs among PFAS. For this, we employed permutation-based significance testing to determine whether some genes were concentration-responsive across multiple PFAS, beyond chance variation expected when comparing thousands of genes. An empirical p -value

was calculated by permutation analysis. Figure 5 shows the top 50 genes that were most frequently concentration-responsive (either up- or down-regulated) to PFAS in each cell type. In this figure, genes are ranked by the frequency of them being identified as a significant CRG in response to PFAS treatment. Almost all the transcripts identified in this analysis were upregulated in response to PFAS, and they were also highly cell type-specific. Only one transcript, *ADAMTS9*, encoding a disintegrin and metalloproteinase with thrombospondin motifs, a gene that is expressed ubiquitously in multiple cell types, was significantly concentration-responsive to PFAS treatment in both iPSC-Hep and iPSC-CM. Genes known to play a role in xenobiotic metabolism or muscle contraction pathways were among the top 50 CRGs in iPSC-Hep or iPSC-CM, respectively. While many of the effected genes were shared among PFAS in each cell type, no gene was shared among more than half of the tested compounds. The gene in highest commonly shared frequency – *CDH6* in iPSC-Hep and *NDUFA10* in iPSC-CM – was significant in concentration-responsive effects in 10 (38%) and 9 (35%) of the tested PFAS, respectively. Still, depending on the gene, anywhere from ~25% to ~40% of the tested PFAS shared common concentration-response effects on gene expression. It is also noteworthy that almost all the shared CRGs in both cell types showed upward concentration-response trends.

Fourth, because we observed that common individual CRGs are known to be involved in the mechanisms of hepato- (Rusyn et al., 2021) and cardiotoxicity (Lind et al., 2021), we aimed to investigate shared pathway-based effects of PFAS and whether these effects are dependent on their chemical structure-based class. For this, we employed the Gene Set Enrichment Analysis approach (Subramanian et al., 2005) to overcome the limitations associated with the utilization of arbitrary thresholds during the selection of significantly differentially expressed genes. This method involves evaluation of cumulative changes in gene expression across entire sets of genes, rather than focusing on specific genes that exceed a predetermined significance threshold. To execute this analysis, we utilized ranked log₂-fold change values obtained by comparing expression levels of each gene at the highest concentration tested with that in vehicle controls. A complete list of gene sets that were significant is included in Supplementary Files 6⁷ and 7⁸, and the corresponding heatmaps using the data are included in Figures S4 and S5⁵. To enhance clarity and interpretability of the outcome of this analysis, an additional data integration step was undertaken to group significantly enriched gene sets from lower-level pathways into common higher-level nodes based on the Reactome pathway hierarchy (Jassal et al., 2020). This organization allowed for a clearer depiction of the common mechanistic events that were affected by tested PFAS. Figure 6 shows that as many as 25 and 18 (out of 26 tested) PFAS exhibited similarity in a number of pathway-level effects in iPSC-Hep and iPSC-CM, respectively. In iPSC-Hep, significantly enriched and upregulated pathways included extracellular matrix organization, translation, RNA and

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⁷ doi:10.14573/altex.2311031s6

⁸ doi:10.14573/altex.2311031s7

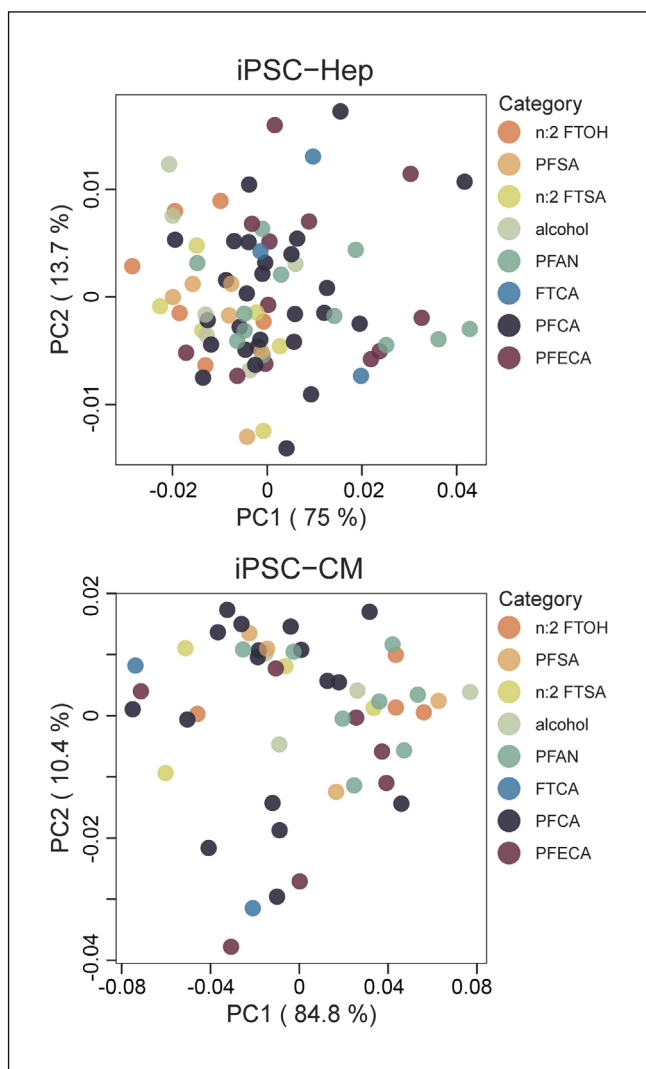


Fig. 3: Principal component analysis (PCA) of normalized gene expression data from iPSC-Hep (top) and iPSC-CM (bottom) treated with PFAS (10 μ M)

In both panels, a dot represents an individual sample, while the colors correspond to different PFAS categories as indicated in the legends. See Tab. 1. for class abbreviations.

general metabolism, and cellular response to stress. Pathways for plasma lipoprotein assembly, remodeling, and clearance were consistently down-regulated in iPSC-Hep by the majority of tested PFAS. In iPSC-CM, all significant pathways were upregulated, and the top effected one was related to muscle contraction, followed by several pathways that are related to energy supply, i.e., events that are required for efficient muscle contraction. We also examined whether chemical structure-based class-specific effects could be discernable at the pathway level in either cell type, using a heatmap to visualize the enriched pathways versus individual PFAS, but found none (data not shown), indicating similarity in transcriptomic effects for PFAS as a class at the pathway level rather than similarity in sub-group.

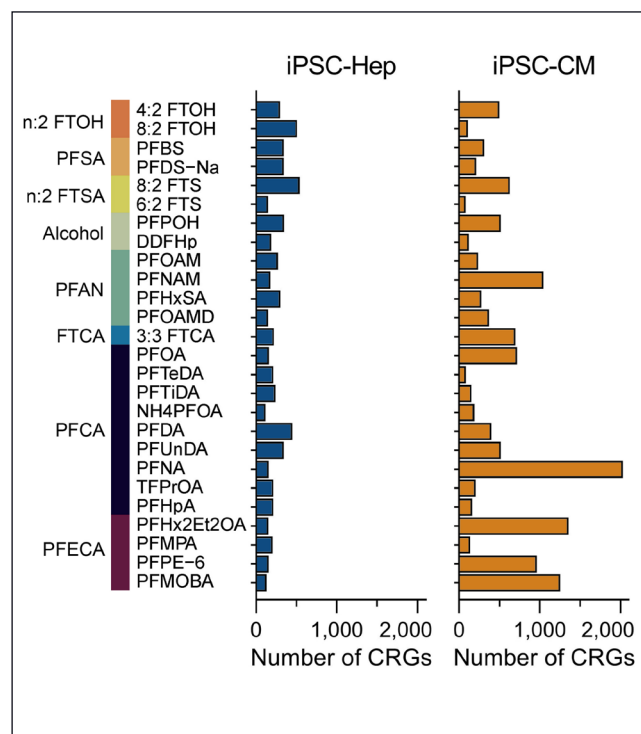


Fig. 4: Bar plots illustrating the number of concentration-responsive genes (CRGs) in iPSC-Hep (displayed as blue bars on the left) and iPSC-CM (shown as orange bars on the right) for each tested PFAS

The PFAS are ordered based on their respective categories. See Tab. 1 for class and individual chemical abbreviations.

Fifth, we calculated transcriptomic PODs (tPODs) at the pathway/gene set level. Recent studies demonstrated the value of using transcriptomic data for dose-response analysis in addition to their traditional use for mechanistic interpretation (Farr and Dunn, 1999; Johnson et al., 2020; LaRocca et al., 2017; Vinken et al., 2017). We used the most sensitive pathway method to derive tPODs, a common approach to dose-response modeling in transcriptomic datasets (NTP, 2018). For each substance, the tPOD is a benchmark dose of the median gene in the most sensitive pathway/gene set effected by PFAS treatment (see Section 2). Figure 7 shows a comparison between tPODs and numbers of CRGs for each PFAS and cell type. A wide range in tPODs was observed among PFAS in each cell type. In iPSC-Hep, only 3 (12%) of 26 tested PFAS had no tPOD, and 23 PFAS had a tPOD < 10 μ M. A significant negative correlation was observed between tPODs and the number of CRGs in iPSC-Hep – as the number of CRGs increased, the tPODs value decreased (correlation coefficient, $r = -0.5$, p -value < 0.01). In iPSC-CM, 10 (38%) of 26 tested PFAS were without tPOD, and 16 PFAS had tPOD < 10 μ M. No correlation was found between tPODs and CRGs. In both cell types, tPODs did not cluster by their chemical structure-based class. No significant correlation was found between the tPODs and either chain length ($\rho = -0.16$ in iPSC-CM; $\rho = 0.003$ in iPSC-Hep) or molecular weight ($\rho = -0.06$ in iPSC-CM; $\rho = -0.05$ in iPSC-Hep) for tested PFAS.

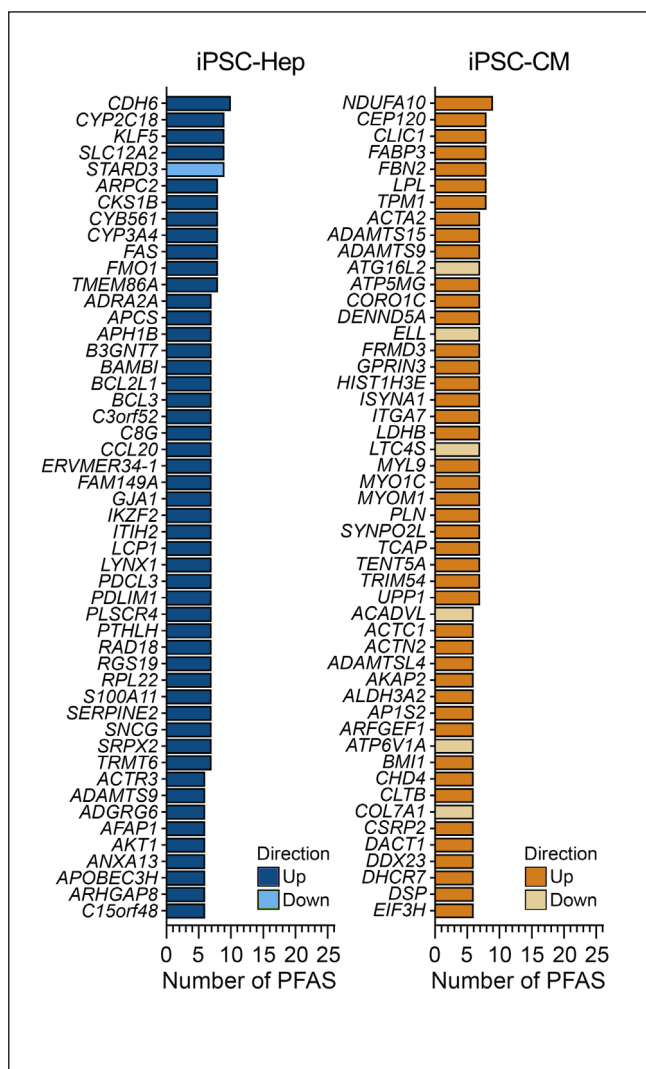


Fig. 5: Bar plots displaying the top 50 common genes that exhibited concentration-responsive effects, ranked by the number of tested PFAS with significant impact (empirical p -value < 0.05 , calculated using permutation) in iPSC-Hep (depicted as blue bars on the left) and iPSC-CM (illustrated as orange bars on the right). Bars with darker shades represent upregulation and lighter shades represent downregulation.

3.3 Risk characterization and prioritization of PFAS using *in vitro* PODs

The analyses presented above showed that the hypothesis that structure-based grouping of PFAS can be substantiated using *in vitro* bioactivity data from both phenotypic and transcriptomic endpoints was not confirmed with the data from the cell types we tested. However, *in vitro* data collected in these studies did show that PFAS elicit both phenotypic and transcriptomic bioactivity. Therefore, we reason that these data may be used for risk-based prioritization of PFAS. One approach is to prioritize PFAS for fur-

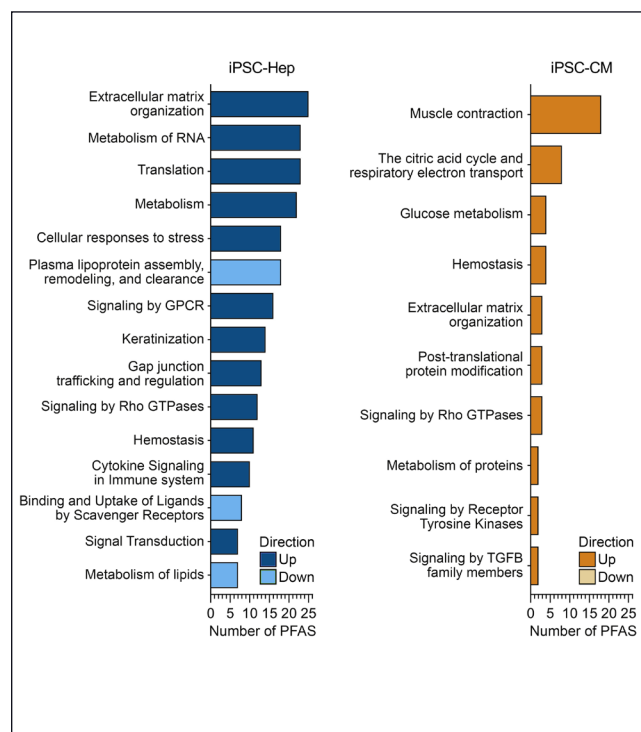


Fig. 6: Bar plots presenting the top groups of Reactome pathways enriched in iPSC-Hep (depicted as blue bars on the left) and iPSC-CM (shown as orange bars on the right), ranked by the number of tested PFAS with significant effects (false discovery $q < 0.1$).

Pathways are color-coded, with darker shades indicating upregulated pathways and lighter shades representing downregulated pathways. The pathways showcased in this figure are summarized based on shared higher-level categories, capturing overarching hierarchical relationships within Reactome pathways. See Supplementary Files 6⁷ and 7⁸ for the full list of enriched pathways and their corresponding upper-level groupings.

ther testing using either phenotypic or transcriptomic PODs alone – the “protective” approach (Woodruff et al., 2023). The other is to combine the “protective” bioactivity-derived PODs and conservative exposure predictions to derive chemical-specific BERs (Paul Friedman et al., 2020). Figure 8 presents both considerations as possible means for prioritization and risk characterization of PFAS. Figure 8A plots the minimum (most protective) pPODs and tPODs for each tested PFAS separately for each cell type. The tPODs were more sensitive (i.e., protective) overall to treatment with PFAS. Specifically, tPODs were lower than pPODs not only on average across all 26 tested PFAS, but also for 25 and 21 compounds in iPSC-Hep and iPSC-CM, respectively.

Next, the overall minimum POD for each substance (either tPOD or pPOD, from either cell type) was used to calculate the BER (Fig. 8B). We used the 95th percentile of the exposure estimates from ExpoCast (Wambaugh et al., 2013), where available, for each PFAS; these were converted to concentrations as detailed in Section 2. Data for both bioactivity and exposure predictions

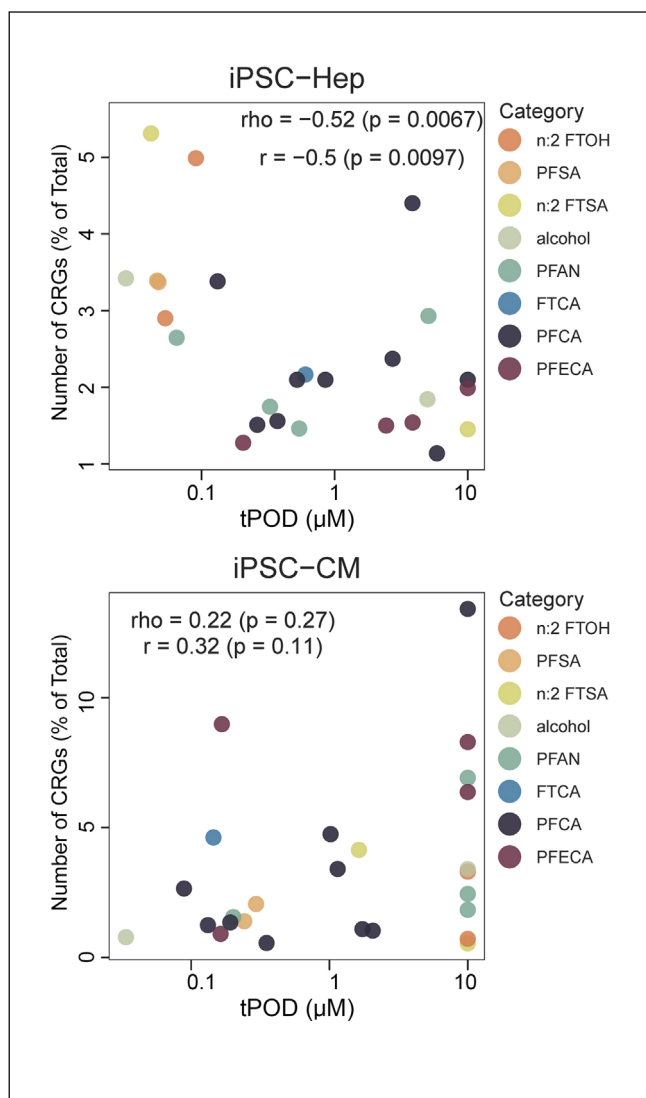


Fig. 7: Correlations between transcriptomic point of departure (tPOD) and the number of concentration-response genes (CRGs, expressed as a fraction of total genes retained after low count removal, as described in Section 2) in iPSC-Hep (top) and iPSC-CM (bottom) treated with PFAS

Both Pearson (*r*) and Spearman (ρ) correlations, and their corresponding *p*-values are displayed. Each data point represents a distinct PFAS, and the colors indicate different categories (as shown in the inset). See Tab. 1 for PFAS class abbreviations.

were available for 18 out of 26 PFAS; therefore, only these substances are included in Figure 8B. Only 3 PFAS exhibited BERs lower than 1 (PFBS, 8:2 FTS and NH4PFOA); most substances ranged between 1 and 100. This suggests that when the most conservative parameters were used for both bioactivity and exposure, several PFAS compounds could be identified as of potentially high concern and may warrant additional studies to refine estimates of both human health hazard and exposure.

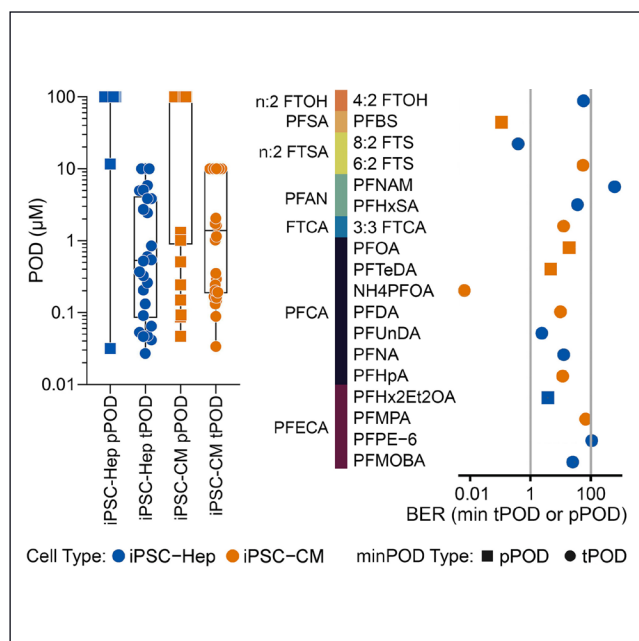


Fig. 8: Comparisons of the points of departure (PODs) derived from phenotypic (pPOD) and transcriptomic (tPOD) data, and the calculated bioactivity-to-exposure ratios (BER) based on the lowest POD for each tested PFAS

(A) POD comparisons. Box-and-whisker plots illustrate the POD values derived from phenotypic data (indicated by solid squares) and transcriptomic data (indicated by solid circles) for iPSC-Hep (blue) and iPSC-CM (orange). The boxes represent the interquartile range, the horizontal line represents the median value, and the whiskers extend to encompass the maximum and minimum values. Values for each tested PFAS are shown as individual symbols. (B) Dot plot displaying the BERs calculated using the lowest (either pPOD or tPOD) values for each tested PFAS with available *in vitro*-to-*in vivo* extrapolation data. The solid dots and solid squares correspond to POD values from phenotypic and transcriptomic data, respectively. The blue and orange colors correspond to iPSC-Hep and iPSC-CM, respectively. The grey lines represent BER at 1 and 100. PFAS are grouped by their chemical structure-based class. See Tab. 1 for class and individual chemical abbreviations.

4 Discussion

Grouping and read-across are widely used approaches, albeit to a varying degree of success, in evaluation of hazards and risks of chemicals that have data gaps (Ball et al., 2016). These approaches are pragmatic means to address the daunting task of evaluating PFAS, an overwhelming majority of which have no data to inform traditional hazard and risk evaluations. One commonly invoked grouping strategy for PFAS involves prioritization based on chemical structure features, which are believed to be most important in determining both toxicokinetic and bioactivity properties (Buck et al., 2011, 2021; Sha et al., 2019; Wang et al., 2017;

OECD, 2018; Patlewicz et al., 2019). In this context, our study aimed to use both transcriptomic and phenotypic data from two human-health relevant (in terms of organs with known or potential PFAS hazardous effects) organotypic (in terms their functionality) *in vitro* cell models to evaluate the association between structural groupings and bioactivity. In addition, regardless of whether structure-based grouping could be substantiated, we evaluated whether concentration-response analysis from both the transcriptomic and phenotypic data, combined with exposure estimates, may offer a sensible approach for risk-based prioritization of PFAS for additional exposure and hazard testing.

4.1 Relationship between PFAS structure-based classes and *in vitro* bioactivity

Several previous studies reported correlations between specific structural features of PFAS and their biological effects *in vitro*. For example, a study that examined effects of 142 PFAS in HepG2 cells (Houck et al., 2021) reported that PPAR activity correlated with physicochemical properties such as the presence of negatively charged structures and acidic hydrogen atoms; they also observed that RXR β activation was associated with linear shapes and carboxylate groups. In another study of 15 PFAS (Zhang et al., 2014) the authors demonstrated that the binding affinity of PPAR γ in HepG2 cells was linked to the chain length. Similarly, a study of 13 PFAS (Amstutz et al., 2022) found that increasing carbon chain length led to higher cytotoxicity in HepG2 cells. PFAS molecular weights were significantly correlated with protein binding association constants and contributed to lipid accumulation (Marques et al., 2022). In addition, a study of 147 PFAS identified a trend between molecular weight and bioactivity related to *in vitro* immunosuppression (Houck et al., 2023), and a study of 160 PFAS discovered that compounds with long carbon chains (8 or more carbons), high carbon to fluorine ratio, or containing a carboxylic acid moiety were more likely to exhibit developmental neurotoxicity *in vitro* (Carstens et al., 2023).

Still, little is known regarding structure-based group similarities in bioactivity even though the latter is a common hypothesis (Buck et al., 2011, 2021; OECD, 2018; Patlewicz et al., 2019). Our study examined potential associations between structural features detailed in previous studies and bioactivity in both iPSC-Hep. While we found few effects in iPSC-Hep, the most noteworthy observation with respect to whole cell bioactivity effects in our study comes from the data in iPSC-CM, a cell type that has not been studied with diverse PFAS previously. Our observation that 8 of 26 tested PFAS had effects on the beat frequency is novel and one that warranted additional mechanistic investigation using transcriptional data. Regardless of the cell type, however, we found little evidence of grouping similarity in the phenotypic effects. Moreover, in contrast to previous findings, our analysis also revealed no significant correlation between PFAS molecular weight or carbon chain lengths and bioactivity. The lack of significant correlations could be due to the number of PFAS used in this study or the cell types used. Further research using larger libraries of PFAS and additional cell types may be needed to confirm potential structure-activity relationships.

4.2 Mechanistic similarity in the transcriptomic effects of PFAS

To further test whether molecular-level events may show evidence of similarity in biological responses to different PFAS, we used gene expression data. Indeed, demonstration of similar mode of action (MOA) has been suggested as a crucial consideration for attaining confident categorization/grouping of PFAS for human health risk assessment (Anderson et al., 2022). Accordingly, we focused on the analysis of gene expression data in several ways, including concentration-response analysis and mechanistic interpretation. Previous studies of diverse PFAS in human hepatocytes or liver-derived cancer cell lines observed accumulation of intracellular lipids for several PFAS at concentrations as low as 0.1 μ M (Marques et al., 2022; Louisse et al., 2023). Similarly, gene expression studies in liver cells suggested that PFAS' structural features may be associated with lipid accumulation-related gene expression effects, including the relative potency of these effects (Reardon et al., 2021; Marques et al., 2022). These studies examined the relationships based on specific structural features, such as aliphatic chain length and molecular weight; they were not based on head group/side chain classification (OECD, 2018; Buck et al., 2011, 2021).

In our study, transcriptomic data in iPSC-derived hepatocytes confirmed the disruption of lipoprotein transport as the most prominent molecular event elicited by many of the tested PFAS. This observation aligns well with previous *in vivo* animal and epidemiological studies that established linkages between PFAS and alterations in serum lipoproteins and liver steatosis (Ho et al., 2022; David et al., 2023; Bijland et al., 2011). Indeed, a number of *in vivo* (Bijland et al., 2011; Curran et al., 2008; Das et al., 2017; Menger et al., 2020; Wan et al., 2012) and *in vitro* (Bjork et al., 2011; Hickey et al., 2009; Naile et al., 2012; Reardon et al., 2021; Rowan-Carroll et al., 2021) studies documented the disruptive impact of PFAS on lipid metabolism in the liver. Furthermore, gene expression data from iPSC-derived hepatocytes also suggested that PFAS may contribute to liver fibrosis, potentially via the up-regulation of TGF- β . This finding supports the reported epidemiological and *in vitro* associations between fibrosis indicators and PFAS exposure (Cheng et al., 2023; Qi et al., 2023). However, our data show that these pathways were neither commonly enriched across all tested PFAS, nor was the significant enrichment associated with the chemical structure-based classes. It remains unclear whether the lack of an effect by some tested compounds was due to the tested concentrations being insufficient (highest concentration examined for gene expression was 10 mM), the choice of a liver cell type, or if some PFAS do not act on these pathways. *In vitro* models often require higher concentrations to demonstrate significant mechanistic results, whether in the case of human primary hepatocytes (Reardon et al., 2021; Rowan-Carroll et al., 2021; Marques et al., 2022; Robarts et al., 2022) or liver cancer cell lines (Solan and Lavado, 2023; Louisse et al., 2023; Robarts et al., 2022). Future research is needed to expand the range of tested PFAS, consider a broader spectrum of concentrations, and include more complex and physiologically-relevant liver model systems (Soldatow et al., 2013).



As we noted previously, published data on the possible associations between PFAS and cardiovascular adverse outcomes is less clear and not as abundant as that for liver. Some epidemiological studies showed the association between PFAS exposure and cardiovascular diseases (Meneguzzi et al., 2021); however, the evidence regarding associations between PFAS and overt cardiovascular disease remains limited and inconsistent (Schillemans et al., 2023). A better understanding of potential underlying molecular events in cardiomyocytes may help explain inconsistencies in epidemiological data; however, studies of heart-derived cells in toxicology are few (Burnett et al., 2021a). Mechanistic omics and laboratory animal studies do indicate the potential for associations of some PFAS with multiple molecular pathways that could contribute to cardiovascular disease, but more information is needed (Schillemans et al., 2023). For example, while some studies suggested that PFAS may affect developmental pathways during cardiomyocyte differentiation (Cheng et al., 2013; Davidsen et al., 2021, 2022; Tang et al., 2017; Zhang et al., 2016), few reports have been published to show effects on adult cells, especially of human origin, and a relatively small number of PFAS has been tested in human cardiomyocytes in culture (Burnett et al., 2021b).

In addition, while iPSC-derived cardiomyocytes are a useful model for cardiotoxicity screening, both in studies of cell function (Burnett et al., 2019, 2021a; Chen et al., 2021; House et al., 2021) and in gene expression studies (House et al., 2022; Tsai et al., 2023; Grimm et al., 2018), gene expression effects from PFAS exposure in cardiomyocytes have not been reported. In this respect, our study provides a unique dataset that shows that PFAS exposure can perturb gene expression in iPSC-CM with patterns that are distinct from those in iPSC-Hep. Specifically, not only did we observe acute effects on the beat frequency, but we also observed that genes and pathways related to muscle contraction and energy supply were significantly affected, also in a concentration-response manner (see below), by several PFAS. These findings suggest that further investigation is needed of PFAS-associated effects in human cardiomyocytes.

4.3 Using transcriptomic data for concentration-response analysis

A number of studies in the past decade showed that the PODs derived from gene expression data in short-term animal studies are correlated with PODs from traditional apical (histopathology or clinical chemistry) endpoints in sub-chronic and chronic animal studies (Thomas et al., 2011; Johnson et al., 2020). More recently, gene expression data have been suggested to be of potential use to set acceptable exposure limits for environmental chemicals – the US EPA has proposed to use transcriptomic reference values (TRV) as a novel EPA Transcriptomic Assessment Product (ETAP) (US EPA, 2023). Therefore, transcriptomic data are not only regarded as useful in mechanistic interpretation, but also for dose-response steps in risk assessment (Nyffeler et al., 2022; Harrill et al., 2021; Tsai et al., 2023; Johnson et al., 2022). Interestingly, a study by (Reardon et al., 2021) examined PFAS-associated *in vitro* tPODs in primary human liver spheroids and demonstrated that they may be more “protective” in comparison to apical PODs

derived from animal studies for most chemicals analyzed, which further demonstrated the value of the use of *in vitro* transcriptomic data in risk assessment.

Accordingly, we aimed to analyze gene expression data in this study with respect to concentration-response. Several previous publications explored gene expression effects of PFAS (Houck et al., 2021; Reardon et al., 2021; Rowan-Carroll et al., 2021; Louisse et al., 2023; Marques et al., 2022; Behr et al., 2020), but few conducted formal transcriptomic benchmark dose analysis and derived points of departure. Two studies used primary human liver spheroids and assessed *in vitro* transcriptomic dose responses of PFAS treatment (Reardon et al., 2021; Rowan-Carroll et al., 2021). Specifically, Reardon et al. (2021) found that PFAS-elicited transcriptomic effects (tPODs) ranged from 0.8 μM to 19.5 μM . Rowan-Carroll et al. (2021) reported that tPODs ranged from 8.2 μM to 53.5 μM . Compared with these two studies, tPODs derived herein were at the lower end (0.03 μM to 10 μM), suggesting that iPSC-Hep could be a more sensitive *in vitro* model for screening-level risk assessments.

In addition to deriving the tPODs, we also sought to explore the relationship between a common tPOD derivation approach – the most sensitive pathway/gene set – that captures molecular pathway/gene set-level bioactivity and the number of CRGs effected by each compound in either cell type. The number of CRGs following chemical exposure has been used as a quantitative indicator of a chemical’s potency to enable comparative analysis across and within chemicals/groups (House et al., 2017, 2022). In addition, CRGs have been proposed to be useful for establishing quantitative thresholds to classify chemicals based on their potential to induce specific outcomes (Ramaiahgari et al., 2019). Our findings indicated that concordance between tPODs and numbers of CRGs was only observed in iPSC-Hep, but not in iPSC-CM. This finding suggests that associations between tPOD-based potency (derived from the most sensitive pathway/gene set) with the overall transcriptomic effect (as determined by the number of CRGs) is cell-type specific. The lack of correlation observed in iPSC-CM may be in part explained by the comparatively smaller evidence base on gene expression effects of chemicals in cardiomyocytes, and thus, a less advanced understanding of the molecular pathways/gene sets that may be impacted (Hudson et al., 2012; Khatri et al., 2012). While *in vivo* and *in vitro* studies have demonstrated comparable tPOD values using different derivation approaches, these tPODs were often derived from few tissues (Farmahin et al., 2017; Thomas et al., 2013) or specific cell types (Reardon et al., 2023). The cell type-specificity observed in this study underscores the necessity for further research to determine what impact certain tPOD derivation approaches, and tissues or cells from which they originate, may have on the sensitivity of these “alternative” PODs, especially as they are proposed for wider use in decision-making (US EPA, 2023).

4.4 Prioritization and risk characterization of PFAS using pPODs and tPODs

The BER approach has been proposed as a high-throughput means of identifying potential risks and guiding the prioritization of substances for higher-tier evaluations (Wetmore et al., 2015; Sipes et



al., 2017; Paul Friedman et al., 2020). Here we applied a highly conservative approach by using the minimum POD for each individual PFAS for the BER derivation, regardless of cell type and assay. We found that both iPSC-Hep and iPSC-CM contributed equally to the lowest PODs, and transcriptomic data contributed 14 of the 18 lowest PODs. These findings underscore the value of both cell models and also emphasize the importance of incorporating both the phenotypic and transcriptomic data for prioritization of PFAS using “protective” *in vitro* data.

We found that the BERs for most PFAS tested in this study ranged from 1 to 100. The overall range of BERs for PFAS in this study is consistent with previous *in vitro* PFAS studies that took a risk-based approach to interpretation of their *in vitro* data. For example, Burnett et al. (2021b) demonstrated relatively narrow margins of exposure (< 100) for most PFAS tested using iPSC-CM phenotypic assays. Rowan-Carroll et al. (2021) also reported a BER for PFOA within one order of magnitude of our study. It should be noted that 3 compounds (PFBS, 8:2 FTS and NH4PFOA) in our study had BERs lower than 1. This apparent indication of a potential high risk was examined further. Both NH4PFOA and PFBS had the highest exposure prediction values after IVIVE transformation (20.5 and 0.43 μM , respectively) among all 26 tested compounds; 8:2 FTS had a very low tPOD (0.04 μM).

We need to acknowledge the limitations with the BER approach for PFAS. First, this study utilized two cell types that represent liver and heart to derive quantitative estimates of bioactivity. While hepatocytes and cardiomyocytes are the primary functional cells in these organs, they may not fully represent complex cell-cell interactions *in vivo*. Therefore, our bioactivity data may not adequately capture organ-level effects. Additional studies in more complex *in vitro* models, as well as cells that represent other tissues, are needed. Second, exposure values used in our study were based on *in silico* predictions rather than empirical data. Third, multiple data sources that we used to inform IVIVE have limitations with respect to the uncertainties in chemical-specific kinetic information for PFAS (Kreutz et al., 2023; Smeltz et al., 2023). Collectively, the uncertainty in BER estimates for PFAS is considerable, and more confident toxicokinetic data is needed for these substances. In addition, despite our best efforts to maximize the number of PFAS for which BERs could be calculated, there were still 8 out of 26 PFAS with insufficient exposure prediction or C_{ss} data for the extrapolation. This underscores the importance of establishing PFAS-specific toxicokinetic and exposure data for large-scale risk prioritization. Still, despite the limitations in the IVIVE process and subsequent BER calculation, the overall workflow herein effectively utilizes existing data and computational tools to enable an informative risk characterization approach that could also be applied for prioritizing PFAS.

5 Conclusion

This study pursued two main objectives, (i) to integrate transcriptomic and phenotypic data from two human *in vitro* cell models to gain insights into structure-effect relationships for PFAS, and (ii) to characterize potential risks of PFAS based on both bioactiv-

ity and predicted exposures. We observed that a number of PFAS had effects on iPSC-CM, supporting limited epidemiological data on the potential for these chemicals to have adverse effects on the heart rhythm. Even though a number of cell type- and substance-specific effects were observed, these effects did not align with the structure-based classification; we found that many of the tested PFAS elicited similar gene expression signatures. These results imply that existing classification proposals may not be suitable for “dimensionality reduction” among PFAS and show that additional research is needed to identify appropriate groupings for read-across. While our data revealed a relatively low risk for almost all tested compounds, even when using most conservative estimates for hazard and exposure, we also acknowledge considerable uncertainties in this approach because of the paucity of relevant toxicokinetic and exposure information. Overall, our results suggest that future considerations of hazards and risks of PFAS may require testing of the individual compounds, rather than representative members of various groups, and that additional studies are needed to inform grouping strategies and risk characterization.

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Conflict of interest

The authors declare no conflicts of interest.

Data availability

Gene expression data and experimental metadata are available in the Gene Expression Omnibus (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE244110>). All supplementary data described in this manuscript are made public.

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