#### A curated collection of human vaccination response signatures 1

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#### 51 Abstract

Recent advances in high-throughput experiments and systems biology approaches 52 have resulted in hundreds of publications identifying "immune signatures". 53 54 Unfortunately, these are often described within text, figures, or tables in a format not 55 amenable to computational processing, thus severely hampering our ability to fully

56 exploit this information. Here we present a data model to represent immune 57 signatures, along with the Human Immunology Project Consortium (HIPC) Dashboard 58 (www.hipc-dashboard.org), a web-enabled application to facilitate signature access 59 and querying. The data model captures the biological response components (e.g., 60 genes, proteins, cell types or metabolites) and metadata describing the context under 61 which the signature was identified using standardized terms from established 62 resources (e.g., HGNC, Protein Ontology, Cell Ontology). We have manually curated 63 a collection of >600 immune signatures from >60 published studies profiling human 64 vaccination responses for the current release. The system will aid in building a 65 broader understanding of the human immune response to stimuli by enabling 66 researchers to easily access and interrogate published immune signatures.

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## 68 Introduction

69 Systems-level profiling of the human immune system has generated important 70 insights into the mechanisms by which humans respond to exposures such as vaccination. These studies, including many conducted through the Human Immune 71 72 Project Consortium (HIPC), have generated hundreds of publications. While 73 repositories exist to promote re-use of primary experimental immunology data generated from these efforts, such as the Gene Expression Omnibus<sup>1</sup> (GEO) and the 74 75 NIAID Division of Allergy, Immunology, and Transplantation (DAIT)-sponsored Immunology Database and Analysis Portal<sup>2</sup> (ImmPort), there is no centralized 76 framework to aggregate and organize the published findings resulting from the 77 78 analysis of this data, and particularly the coherent sets of biomarkers, termed here 79 "signatures". Additionally, such signatures are not published in a consistent format 80 between publications and may be presented as text, tables, or images. This 81 heterogeneity presents a barrier to comparative analyses since identifying published 82 signatures, for example of a vaccine response, requires extensive manual curation of 83 the literature that must be repeated by investigators each time they wish to interpret a 84 set of results. Here, we propose a model to standardize the representation of these 85 published findings and present the Human Immunology Project Consortium (HIPC) 86 Dashboard—a searchable interface to query curated signatures from the corpus of 87 human immunology literature.

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89 We define a 'signature' as the information required to specify a published result. This 90 includes alterations in the levels of a set of one or more response component(s), i.e., 91 biological entities such as genes or cell types, that are defined by a particular 92 comparison in the context of an immune exposure. The signature also includes 93 contextual information (termed metadata) such as the conditions and circumstances 94 under which the signature was identified, the tissues or cells that were assayed, as 95 well as clinical data such as demographic information about the groups that were included in the analysis. As a motivating example, a study by Bucasas et al.<sup>3</sup> 96 97 identified a set of genes that are up-regulated in individuals with higher antibody 98 responses (comparison) after vaccination with the 2008-2009 trivalent influenza 99 vaccine (exposure) in an adult cohort. The expression of genes STAT1, IRF9, SPI1, 100 CD74, HLA-E, and TNFSF13B one day after influenza vaccination was predictive of 101 greater antibody responses. In the paper, these results were represented in a table, 102 though similar findings often appear as text or within figures. Without standardization, 103 such findings are not easily accessible to the wider scientific community for further 104 analysis.

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Several existing resources define pathway and gene module signatures through re analysis of raw data, but few capture the original findings published with these data or
 are specifically geared towards human immunology research. Among these
 resources are the OMics Compendia Commons (OMiCC)<sup>4</sup>, EnrichR<sup>5,6</sup>, the integrative
 Library of Integrated Network-based Cellular Signatures (iLINCS)<sup>7</sup>, the Molecular

Signatures Database (MSigDB)<sup>8,9</sup>, and VaximmutorDB<sup>10</sup>. OMiCC crowdsources 111 112 annotations for gene expression data to be used in re-analysis and novel signature 113 generation. EnrichR and iLINCS offer biological annotations built from data re-114 analyzed en masse, but similarly do not capture published findings. MSigDB does 115 include manually curated gene signatures along with those derived from data re-116 analysis, albeit with fewer contextual details than captured for the HIPC Dashboard. 117 VaximmutorDB captures published gene expression and proteomic signatures but not 118 cell-type frequency signatures, and signatures from this database are not yet 119 downloadable in a machine-readable format.

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121 To improve access and to promote reuse of published signatures, we designed a 122 data model that standardizes the content and context of published immune 123 signatures. Our initial curation efforts have focused on gene expression and cell-type 124 frequency/activation signatures of human vaccine responses, but this framework is 125 extensible to other domains such as response to infection. We captured what is 126 changing, (e.g. groups of genes), how that response component changed (e.g. up- or 127 down-regulation), where this change was observed (e.g. in sorted CD8+ T cells from 128 adults), and the comparison that was performed (e.g. individuals with high vs. low 129 antibody titers post-vaccination). We then manually curated signatures from 130 publications both within and outside of HIPC that described changes in gene 131 expression, cell-type frequencies, or cell activation state in response to vaccination. 132 To disseminate these immune signatures, we developed the HIPC Dashboard 133 (www.hipc-dashboard.org), a web-accessible, user-friendly interface to enable 134 signature searching and browsing, and to facilitate rapid comparative analyses. The 135 design of the HIPC Dashboard is based on a similar infrastructure we developed 136 previously for the Cancer Target Discovery and Development network, the CTD<sup>2</sup> 137 Dashboard<sup>11</sup>, and leverages the same underlying ontological framework for the 138 standardized representation of research findings as well as the emphasis on the 139 consistent, curation-mediated use of controlled vocabularies for linking findings 140 reported in different publications.

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### 142 **Results**

### 143 A Data Model for Immune Signatures of Vaccination

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145 We developed a data model that captures, in a detailed and consistent format, the 146 essential information embedded in published immune signatures of vaccination for 147 dissemination through the HIPC Dashboard (Table 1). Key elements of this data 148 model (e.g., genes, vaccines, etc.) are specified using controlled vocabularies, thus 149 making immune signatures of vaccination amenable to data mining and promoting 150 compatibility with projects both within and outside of HIPC. A signature as defined in 151 this model encapsulates both a change in the behavior or abundance of a biological 152 response component as well as the metadata describing the context under which the 153 signature is identified, including (1) the tissue in which the signature was observed, 154 (2) the immune exposure and timing underlying the observed comparison, and (3) 155 clinical details of the cohort from which tissue samples were taken, including age 156 (Figure 1). The model accommodates many types of biological response 157 components (gene, protein, metabolite, pathway, and cell type (e.g. subsets of blood 158 cells). We focused on gene expression and cell type signatures of vaccination, but 159 the data model and HIPC Dashboard infrastructure are flexible and can be easily 160 expanded to accommodate arbitrary signature types.

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162 To facilitate data mining and comparative analyses between different conditions 163 including vaccine types, and to afford consistency between this database and other 164 projects using the same controlled vocabulary terms, standardized terms and 165 ontology links were used for as many biological response components, immune

exposures, and demographic fields as possible. Gene and cell type response components were standardized to the HGNC<sup>12</sup> (as provided through the NCBI) and Cell Ontology<sup>13,14</sup> (CL) respectively to enable comparisons across publications using different naming conventions. Cell types from the Cell Ontology were further differentiated using protein marker terms drawn from Protein Ontology<sup>15</sup> (PRO), where possible (e.g. IFNG+ T cells). This same naming convention is used to describe the tissue in which the signature was observed<sup>16</sup>.

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To annotate the immune challenges driving each signature, we utilized the Immune 174 Exposure model<sup>17</sup>, which provides a standardized description of a broad range of 175 176 potential and actual exposures to different immunological agents (e.g., vaccination, 177 laboratory confirmed infection, living in an endemic area, etc.). Immune exposures 178 are broken down into Exposure Process, Exposure Material, Disease Name, and 179 Disease Stage. Each of these components is modeled using standardized ontology 180 terminology. Within the data model for the HIPC Dashboard, Exposure Materials such as vaccines are captured using terms in the Vaccine Ontology<sup>18</sup> (VO), which further 181 link to target pathogens and strains using the NCBI Taxonomy<sup>19,20</sup>. While these 182 ontology choices reflect our initial focus on vaccination, the data model can 183 184 accommodate other exposure processes beyond vaccination, with links to 185 appropriate ontologies. Integration of the Immune Exposure model in the HIPC 186 Dashboard data model promotes interoperability with other projects that have adopted its use, both within and outside of HIPC, including data repositories such as 187 ImmPort and the Immune Epitope Database<sup>21</sup>. 188

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190 Cohort information that is important for interpreting signatures is also captured. 191 Cohort descriptors can vary widely between studies and can include, for example, 192 sex, antibody response titers, geographic location, health status, vaccination or 193 infection history, etc. This information is currently recorded as unstructured text to 194 maintain flexibility. Cohort age range is standardized separately by storing minimum 195 and maximum ages along with their units. Additional fields describe the particular 196 perturbations that drive the changes to the biological response components. The 197 "comparison" field describes the cohort groups whose differential response under the 198 perturbation is measured. Examples of comparison groups include measurements 199 taken at two different time points (e.g. day1 vs. day 0), correlation with antibody 200 response, differing antibody response outcomes (e.g. high vs. low responders), or 201 comparisons across different demographic parameters such as age or sex (e.g. 202 younger vs. older, female vs. male). The "response behavior" field captures the 203 directionality of the differential response (e.g. up or down, positively- or negatively-204 correlated) under the specified comparison. Fields for which a formal controlled 205 vocabulary was not used, such as cohort descriptions and the comparison, are stored 206 as free text.

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Finally, each signature is tagged with a Pubmed ID (PMID) and publication year field,
 to connect observations to their literature sources.

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Manual Curation of Published Signatures

213 A defined Pubmed search strategy was used (see Methods) to assemble an initial list 214 of publications comprising studies that involved a systems-level profiling of 215 measurable changes before and after vaccination in human subjects. The 216 publications were culled for signatures reporting statistically significant changes in 217 gene expression, cell-type frequency or cell activation state induced by an immune 218 exposure when comparing groups with different features (such as high vs low 219 responders as defined by antibody titers). We focused on components that also 220 included information about response behavior (e.g. up- or down-regulated). In total,

665 immune signatures were manually curated from 69 published studies. After
standardization and quality control (see Methods), these curated gene and cell type
signatures included 13,812 unique genes, 152 unique cell types (including protein
markers and additional type-modifiers), and 44 pathogens across 56 vaccines (Table
Table 3 illustrates a typical gene-expression type signature after tissue, gene
symbol and pathogen standardization.

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#### 228 The HIPC Signatures Dashboard

229 The data model provides a means for representing immune signatures in a 230 structured, standardized, machine-readable manner while the curation process 231 enables the cross-referencing of signatures from different publications based on their 232 shared response components, by enforcing the consistent use of controlled vocabularies for codifying these components (e.g., genes, cell types, tissues, 233 234 vaccines, and pathogen strains). These capabilities come together in the "HIPC 235 Dashboard" (http://hipc-dashboard.org), a web application developed to enable dissemination of the curated set of immune response signatures. The HIPC 236 237 Dashboard allows signature browsing, as well as searching for one or multiple 238 response components (using the corresponding controlled vocabulary terms and their synonyms), to retrieve all immune signatures involving the query response 239 240 component(s) across all curated publications.

241 The central viewable element of the HIPC Dashboard is the "Observation Summary", 242 a human-readable description of the information captured in an immune signature. 243 Observation summaries are constructed "on the fly", using template text devised as 244 part of the curation process. The template has placeholders for the various elements 245 of an immune signature, including the response component (gene or cell type) and 246 the response behavior type (up/down or correlation). When a specific signature is 247 selected in the process of browsing or searching the Dashboard, the observation 248 summary for that signature is instantiated by replacing the template placeholders with 249 the relevant values from that signature. For example, a joint search on the terms 250 "CD4" and "Zostavax" yields about 35 observation summaries. One of these is related 251 to a change in cell-type frequency:

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In peripheral blood mononuclear cell, CD4-positive, alpha-beta memory T cell & CD38+, HLA-DR+, VZV tetramer+ frequency was up at 14 days from time of vaccination for the comparison 14d vs 0d in cohort 50-75 yo after exposure to Zostavax targeting Human alphaherpesvirus 3 (details\_»)

Here, the "&" sign separates the Cell Ontology cell type from Protein Ontology surface and other markers. In a second example, a joint search on the terms "CXCL10" and "BCG" yields 6 observation summaries, one of which reports a correlation of gene expression (at 1 day post-vaccination) to an ELISpot result at 28 days post-vaccination:

In blood, CXCL10 gene expression at 1 day from time of vaccination was positively
 correlated with IFN-gamma ELISpot spot forming cell 28d in cohort 4-6 mo
 subgroup BCG-primed after exposure to MVA85A targeting Mycobacterium
 tuberculosis variant bovis BCG (details »)

In both cases, placeholders in the observation summary template have been replaced by controlled terms for the response components and ontology-linked metadata (blue, hyperlinked text) and by free-text metadata describing informative experiment details (black, bold text). Following the hyperlink for a controlled term leads to a dedicated page for the corresponding biological entity, providing additional 274 details (including links to relevant external annotation sources, e.g., Entrez, 275 GeneCards and UniProt for genes) as well as a listing of all the immune signatures 276 stored in the Dashboard that involve that entity (Figure 2A). Further, the "details" link 277 at the end of each observation summary points to an "Observation" page (Figure 2B) 278 containing detailed information about the corresponding immune signature, including 279 a full listing of all its available metadata. This includes, for example, structured text for 280 values such as age group and days post-immunization, and links to download the full 281 signature source data (including all metadata) in tab-delimited form. Additionally, 282 each observation includes a link to a file containing the complete set of response 283 components from which it was derived, e.g. the full list of genes or cell types.

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#### 285 Discussion

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287 Users of the HIPC Dashboard can easily search and examine hundreds of immune 288 signatures related to human vaccination responses. Consolidating these publications, 289 standardizing their findings in a database, and disseminating them through the 290 Dashboard interface allows for rapid comparative analyses and re-use of published 291 findings. This is particularly important for identifying commonalities across studies 292 that may reflect shared mechanisms. The HIPC Dashboard can offer broad insights 293 into the mechanisms by which our immune systems respond to vaccination and will 294 be of great value to the vaccine research community. Although the HIPC Dashboard 295 is not designed as an analysis engine, all signatures are made available for download 296 so that users may perform more sophisticated and targeted downstream analyses.

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298 Among data resources dedicated to the collection of vaccination signatures, the HIPC 299 Dashboard is nearly unique in its emphasis on manual curation of published 300 literature. To the best of our knowledge, only MSigDB and VaximmutorDB maintain 301 signatures curated from publications. MSigDB provides minimally redundant gene 302 sets for enrichment analyses, but unlike the HIPC Dashboard does not attempt to 303 capture the full biological context of published results. A reduced set of our curations 304 has recently been made available for gene set enrichment analyses through MSigDB 305 under the C7 VAX gene sets. VaximmutorDB provides access to a collection of 306 immune factors (genes/proteins) that change in response to vaccination against 46 307 pathogens. Compared to VaximmutorDB, the HIPC Dashboard offers several 308 advantages, including: (i) a wider breadth in the types of response components and 309 immune changes that are captured, (ii) improved browsing functionality that facilitates 310 comparisons of immune changes across studies and vaccines, and (iii) the ability to 311 download signature data.

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313 As of the date of this publication, 152 unique cell types and 13,812 distinct genes 314 have been collected in the HIPC Dashboard; this large number of published results 315 allows users to quickly examine the role of particular biological response 316 components, such as individual genes or cell types, across studies. Most of the 317 currently curated gene signatures have fewer than 50 genes, with a range of 1 to 318 2,036 (Figure 3A), while most cell-type frequency or cell activation state signatures 319 have only a single cell type, with a range up to 9 (Figure 3B). These signatures 320 represent findings from 16 tissues and tissue extracts, including blood, PBMCs, T 321 cells, B cells, monocytes, and NK cells. Nearly 250 entries describe changes over 322 time, more than 75 capture antibody response-associated signatures, and several 323 others come from studies that report effects of age and T cell responses.

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The frequency with which cell types and genes are reported in the Dashboard offers insights about key players of the human immune response to vaccination. The most commonly reported genes are STAT1, a key mediator of immune response activated 328 by cytokines and interferons; GBP1, an interferon induced gene involved in innate 329 immunity; IFI44L, a paralog of Interferon Stimulated Protein 44, and SERPING1, a 330 complement cascade protein (Figure 3C). The most common cell types across pathogens and comparisons are NK cells, CD4+ T cells, and CD8+ T cells. (Figure 331 332 **3D).** We searched the HIPC Dashboard data for genes with vaccination signatures 333 across six or more pathogens and found a set of 36 associated genes across 12 334 target pathogens (Figure 3E). Many are interferon stimulated genes, Toll-like 335 receptors, or members of the complement cascade, potentially reflecting a common 336 transcriptional program in response to many different vaccinations.

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338 By design, our current implementation captures vaccination signatures as they are 339 reported in the literature, but it does not include related methodological or statistical 340 information regarding signature discovery (e.g. p-value cut-off) or provide analytical 341 tools that can be applied to the curated signatures. Test statistics are not usually 342 comparable across study designs, and we believe this information may give users a 343 false sense that some signatures are more statistically reliable than others. We 344 instead defer to the judgement of each study's authors and their peer reviewers, and 345 capture signatures as they were reported in each publication. We also caution that 346 bias regarding the number of times particular genes or cell types were investigated 347 might skew relationships in the Dashboard, thus precluding certain types of analyses. 348 As a result, high level analytical tools have not been integrated into the Dashboard, 349 although all of the signatures with full metadata can be downloaded to enable the use 350 of third-party tools. Despite this, we believe the signatures available in the HIPC 351 Dashboard will allow the research community to quickly query the literature and 352 provide valuable comparisons and context for their own experimental results.

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354 The number of genes and cell types captured reflects publications curated through 355 January 2021, but we anticipate the HIPC Dashboard will undergo regular updates to 356 accommodate new findings and additional domains of interest. The current 357 implementation includes gene and cell type response components, as these 358 represent the most commonly published signature types, but it will be valuable to also 359 curate other response components, such as pathways, proteins, and metabolites. 360 Additionally, we recognize that researchers may wish to compare a vaccine response 361 against a particular pathogen to its corresponding disease response; it is easy to see 362 how future iterations could expand the existing vaccine signature framework to 363 capture signatures of infection. Based on our experience, we expanded the data 364 model to include figure numbers or supplementary file annotations within publications 365 as this can greatly simplify quality control during manual curation. We have provided 366 links in the Dashboard to original sources wherever possible. We are also keen on 367 exploring advancements in text-mining and artificial intelligence (AI), to assess how 368 they can assist in automating signature identification and coding. To that end, the 369 immune signatures in the HIPC Dashboard can be used as a data source for 370 training/testing such AI solutions in the future.

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372 In summary, we present the HIPC Dashboard (hipc-dashboard.org) to provide the 373 vaccine research community with easy access to hundreds of published human 374 systems vaccinology signatures. This resource will allow researchers to rapidly 375 compare their own experimental results against existing findings that may otherwise 376 be difficult to locate in the literature. This resource encourages the re-use of 377 published results for advancing our understanding of human vaccine responses and 378 provides a framework that can be extended to capture signatures from other types of 379 immune exposures.

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### 381 Methods

382 Manual Curation

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384 The initial list of publications to curate into the HIPC Dashboard were derived from a 385 PubMed search of papers matching the terms "Vaccine [AND] Signatures" or "Vaccine 386 [AND] Gene expression". Publications were further filtered to meet a set of inclusion 387 criteria: (i) study involved human subjects, (ii) provided a comparison of a measurable 388 change or correlation before and after vaccination (or challenge), and (iii) were 389 reported as statistically significant. Signatures were excluded if they were missing 390 directionality, or if they were derived from datasets external to the publication, to 391 avoid redundancy. Two data curators manually collected a standard set of 392 information from each study according to the designed data model (see Figure 1) 393 and recorded it into a spreadsheet. Each signature was entered by one curator, and 394 subsequently double-checked by the second curator. Table 3 shows a representative 395 portion of the standardized information captured for each signature (12 data fields out 396 of a total of 25).

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398 Assays in the curated publications included gene expression analysis and measured 399 changes in cell-type frequency and cell activation state. Each publication could give 400 rise to any number and type of individual signatures. The signature content is 401 centered on a list of biological response components (genes or cell types) that had a 402 statistically significant change in the assay. These were designated as "response 403 components" to capture different types of entities in a single standardized template 404 column. For example, for gene expression assays, this was often a list of differentially 405 expressed genes.

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407 For genes, an initial manual curation process was applied to make a first pass at 408 symbol standardization and detect any mistakes in copying. Gene names, symbols and or IDs (which may include HGNC symbols, Entrez IDs, Ensembl IDs etc.) were 409 searched in turn against Panther, (www.pantherdb.org)<sup>22</sup>, using the "Functional 410 classification viewed in gene list" search, followed if needed by searches against 411 UniProt (www.uniprot.org)<sup>23</sup> and NCBI (www.ncbi.nlm.nih.gov/search) until either a 412 413 match or updated symbol was found; in the case of no match, the original 414 representation was left unchanged. Any IDs that matched entries that were 415 deprecated or defined as pseudogenes were removed from the curation.

- 416
- 417 Data Standardization
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The manually curated data required further steps to match terms encountered to their appropriate ontology representations. A number of the translations described below were orchestrated using an R script, which generated files ready for loading into the HIPC Dashboard.

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424 Gene symbols - Outdated gene symbols and known aliases were translated to their 425 current NCBI representation, which is the HGNC symbol in all but one case. The first 426 pass of conversion used the function alias2SymbolUsingNCBI() from the Bioconductor limma package<sup>24</sup> with the most recent available gene annotation file. 427 428 This function returns either an exactly matching official symbol, or if none, the alias with the lowest EntrezID. We followed this by a second R package, HGNChelper<sup>25</sup>, 429 430 which was able to resolve additional unmatched gene symbols to valid NCBI 431 symbols. Genbank accession numbers were converted to gene symbols where 432 possible using the org.Hs.egACCNUM2EG translation table which is part of the Bioconductor org.Hs.eg.db package<sup>26</sup>. Selected symbols still not matching NCBI 433 434 names were investigated and corrected manually where possible after checking the 435 original publications for context or for errors in transcription. Symbols for which no 436 valid NCBI gene symbol was found, e.g. some pseudogenes, antisense, or 437 uncharacterized genes, are not included in the HIPC Dashboard proper, since a

requirement of the Dashboard framework is that all gene symbols must appear in the
controlled vocabulary (NCBI/HGNC). However, these symbols are included in
downloadable complete gene lists for each signature.

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442 *Cell types* - Cell types as response components were first curated from the 443 publications as published using a combination of cell type terms and additional 444 descriptive terms, such as protein marker expression. This information was then 445 mapped to a combination of Cell Ontology and Protein Ontology terms, according to a published model<sup>16</sup>. Note that cell types can appear in two different contexts, either 446 as response components themselves, or as the cell type isolated for gene expression 447 448 experiments. In some cases, additional information was provided which could not be mapped to an ontology term<sup>16</sup>. This type of information related to a wide variety of cell 449 450 identification techniques and included the use of additional stains such as viability 451 dyes or tetramer staining. For each set of terms, an entry was created in a lookup 452 table by assigning (1) a parent cell type from the cell ontology, (2) mapping additional 453 protein marker terms to the protein ontology, and then (3) separately retaining as free 454 text descriptors not mapped to an ontology entry, such as tetramer specificity. Thus, 455 the original entry was mapped to up to three descriptor columns, which can be 456 combined as needed for display purposes. For a full, translated cell type, the 457 displayed format is the cell ontology name followed by, if there are additional terms, 458 the "&" symbol, followed by any PRO terms and then any free text.

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Vaccines - Vaccine names collected from the literature were manually mapped to the
most specific vaccine ontology term available. If a specific vaccine could not be found
in VO, new terms were requested. Some examples of terms we included were
VO\_0004899 (2012-2013 seasonal trivalent inactivated influenza vaccine),
VO\_0003961 (ChAd63-KH vaccine, *Leishmania donovani*), VO\_0004890 (gH1-Qbeta
vaccine, novel pandemic-influenza), and VO\_0004891 (CN54gp140□+□GLA, HIV-1).

467 Pathogens - The viral, bacterial or protozoan pathogens targeted by each vaccine are 468 represented with terms from the NCBI Taxonomy. For the case of influenza 469 vaccines, a table was created mapping vaccines by year of administration and type 470 (e.g. trivalent or quadrivalent) to their seasonal viral components, unless otherwise 471 indicated in the publication (e.g. for monovalent or specialized vaccines such as 472 Pandemrix). For the few cases where the exact viral strain was not present in the 473 NCBI Taxonomy, the closest more general term in the hierarchy was used. This 474 mapping table was used to substitute in the actual viral pathogen names. 475

## 476 Data Availability

477 Curated signatures are available on the HIPC Dashboard website (http://hipc-478 dashboard.org) and Github (see Code Availability for details). Files listing all 479 response components for a signature can be downloaded from within individual 480 observations in the Dashboard. The complete set of signature data can be downloaded from the GitHub repository at https://github.com/floratos-lab/hipc-481 482 dashboard-pipeline. This repository contains copies of (1) the original curated data 483 sheets, (2) the response components in individual files, one per signature, (3) the response components in the Broad GMT format<sup>8</sup>, and (4) the actual tab-delimited 484 485 Dashboard load files, in which the complete signature data is fully denormalized into 486 an easy-to-parse format. Further details about the file formats are available on the 487 GitHub project page. R session information for the Dashboard signature pre-488 processing pipeline is available on GitHub.

#### 489 Code Availability

Source curated data and mapping files (cell types, vaccine components, the NCBI 490 491 gene file used, etc.), as well as the R code for the processing pipeline used to create 492 the Dashboard submission files. are available on GitHub at 493 https://github.com/floratos-lab/hipc-dashboard-pipeline. The data in this paper 494 corresponds to pipeline and data version 1.2.1 in the pipeline GitHub repository. 495 Code for the HIPC Dashboard web interface is available on GitHub at 496 https://github.com/floratos-lab/hipc-signature.

- Supported web browsers The HIPC Dashboard has been tested on recent versions
   of
- Chrome (Version 93.0.4577.63 (Official Build) (64-bit))
- Firefox (Version 92.0 (64-bit))
- Edge (Version 93.0.961.38 (Official build) (64-bit))

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## 515 **Competing interests**

516 S.H.K. receives consulting fees from Northrop Grumman and Peraton.

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## 518 **Figure Legends**

**Figure 1.** Overview of the manual curation process for extracting immune signatures from relevant publications into the HIPC signatures database and a web-accessible HIPC Dashboard. The middle panel highlights the various fields that are captured for a given immune signature, with examples provided in red font. Key fields are standardized using existing ontologies or pre-defined criteria in order to capture a wide array of signatures.

525

526 Figure 2. HIPC Dashboard web interface. A. Subject page for cell type "CD4-527 positive, alpha-beta T cell" showing a link-out to the Cell Ontology, the filtering box to 528 further narrow the displayed observations, and the first two observation summaries 529 ("Related observations"). B. Partial view of a details page for a CD4-positive, alpha-530 beta T cell observation. For each controlled term, its name, plus its class, role, and 531 description are shown. Linked pages list details from the relevant ontology and list all 532 observations containing that term. The class equates to its controlled vocabulary 533 type; values are cell subset, gene, pathogen, and vaccine. Roles are used to further 534 differentiate how each term, whether controlled or standardized, is being used.

Among the classes in the HIPC Dashboard, only the class "cell subset" has more than one role, these being "tissue" and "cell\_biomarker". Full metadata, not shown here, is contained in the table labeled "Evidence" at the bottom.

538

539 Figure 3. Summarization of HIPC Dashboard contents. Signature size distributions 540 showing the number of response components across A. gene and B. cell type 541 signatures. C. Word cloud of the top 50 most frequent gene symbols and D. top 10 542 most frequent cell types, where size corresponds to the total number of observations 543 in the Dashboard. E. Heatmap of recurring genes across vaccines targeting different 544 pathogens. Temporally associated genes in adult whole blood or PBMCs were 545 filtered to those with signatures for six or more pathogens. Color indicates up (red) or 546 down (blue) regulation. Genes with opposing directions in multiple studies were 547 marked 'trends up' or 'trends down' according to the most common direction (or 548 marked 'no consensus' for perfect ties).

549

## 550 Tables

551

**Table 1.** Data model for capturing immune signatures. Genes and cell types are captured as response components, with terms standardized against NCBI/HGNC or CL+PRO, respectively. Exposures are captured and standardized against VO terms and NCBI Taxonomy IDs. Metadata includes observed tissue, study timing, cohort descriptors, and age characteristics. \*fields in the Immune Exposure model<sup>17</sup>

557

558 Table 2. Dashboard summary statistics for gene and cell-type signatures. "Joint" 559 refers to the union of the two signature types, as they overlap in the various 560 components. "Total Response Components" lists the number of genes or parent cell 561 types from the Cell Ontology (CL) across all signatures. When additional cell-type markers are included, e.g. from the Protein Ontology (PRO), there are 152 unique 562 "Response Components per 563 cell types represented among the signatures. 564 Signature" shows the range of the number of response components found in 565 individual signatures.

566

Table 3. Key fields in the immune signature data model for a gene expression
 signature. This signature reports positive correlation between gene expression and a
 computed titer response index (TRI).

570

571

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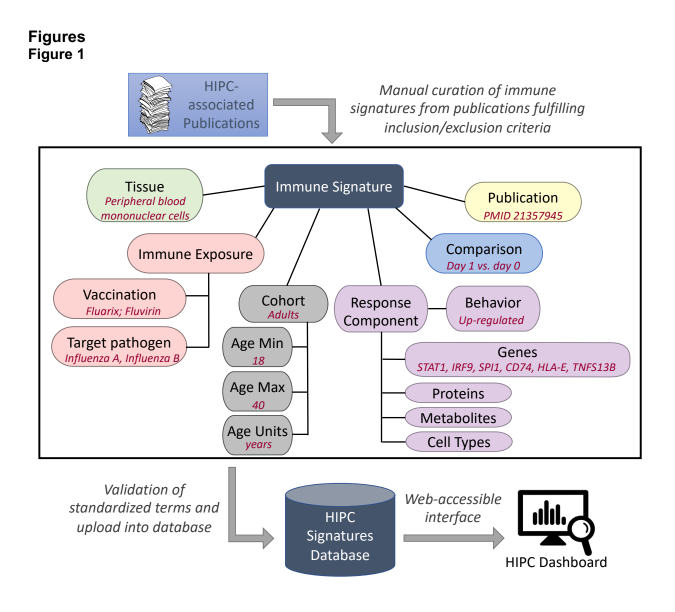
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631

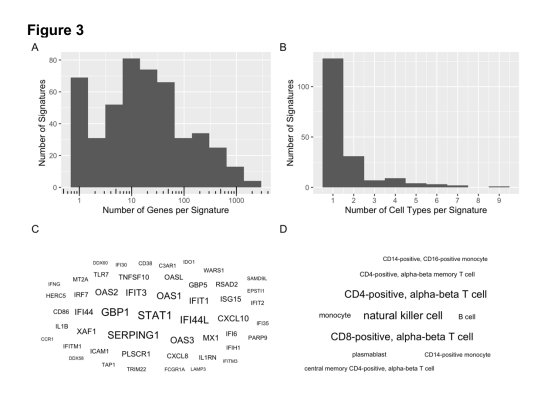


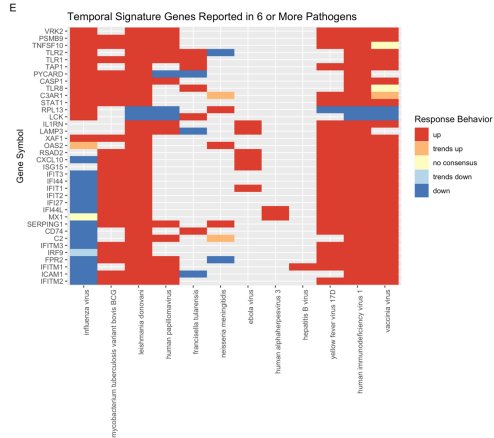
# Figure 2

A CD4-positive, alpha-beta T cell							
Cell Ontology ID	CL_0000624	Cell Subset					
Definition	A mature alpha-beta T cell that expresses an alpha-beta T cell receptor and the CD4 coreceptor.						
Comment							
Broad Synonyms							
Exact Synonyms	CD4-positive, sipha-beta T lymphocyte     CD4-positive, sipha-beta T-cell     CD4-positive, sipha-beta T-lymphocyte						
Related Synonyms							
References	Cell Ontology: CL_0000624 EMBL-EBI OLS: CL_0000624						
Related observations for the role of cell_biomarker							

Showing 1 to 10 of 64 entries Export as Spreadsheet Filter this Table:						
Publication Date v Observation Summary				PMID	÷	
Mar 16, 2019	Mar 16, 2019 In peripheral blood mononuclear cell, CD4-positive, alpha-beta T cell frequency was up at 0 days from time of vaccination for comparison females vs males in cohort 50-74 yo after exposure to Fluarix targeting Influenza A virus (A/California/7/2009(H1N1)), Influenza A virus (A/Perth/16/2009(H3N2)) and/or Influenza B virus (B/Brisbane/60/2008) (details »)				30873150	
Sep 25, 2018		In blood, CD4-positive, alpha-beta T cell & IFNG+, VZV-specific frequency was up at 72 hours from time of vaccination for comparison 72hrs post-vaccination vs pre-vaccination in cohort 70-93 yo after exposure to Zostavax targeting Human alphaherpesvirus 3 (details »)			30247603	

В	B							
Obs	Observation III (1997)							
	In blood, CD4-positive, alpha-beta T cell & IFNG+, VZV-specific frequency was up at 72 hours from time of vaccination for comparison 72hrs post- vaccination vs pre-vaccination in cohort 70-93 yo after exposure to Zostavax largeting Human alphaherpesvirus 3							
	Name	Class	Role	Description	~			
6	CD4-positive, alpha-beta T cell	CellSubset	cell_biomarker	response component				
6	blood	CellSubset	tissue	tissue type				
Q	Human alphaherpesvirus 3	Pathogen	pathogen	target pathogen				
Annet	Sostavax	Vaccine	vaccine	exposure material				
Study (show details)								
Evidence								
Showing	1 to 35 of 35 entries			Filter this Table	e			
÷	Type A Description Details							





Pathogen

Table 1

Elements	Content Definition	Field value Example	Ontology	Data Type	Content Format
Response Component	The biological entity being observed: either a specific gene or cell subset	CKS1B	NCBI/HGNC for genes, Cell Ontology for cell types, PRO for proteins)	String	Controlled
Response Component Type	Type of response agent, e.g. gene, cell subset	gene		String	Fixed List
Tissue Type	The type of cells analyzed, e.g. whole blood, gated cells	whole blood (UBERON:0000178)	Blood + PRO/CL	String	Controlled
Exposure Process Type * Exposure Material * Disease Name *	Category of immune exposure Eg, the type of vaccine administered The condition being observed	Vaccination VO:0001176 None	VO DO	String String String	Fixed List Controlled Controlled
Disease Stage *	Stage of infection (e.g. acute, chronic, post, etc.)	None	OGMS	String	Controlled
Target Pathogen	The pathogenic organism (e.g. virus, bacterium, prion, fungus) being studied	Influenza A, Influenza B	NCBI Taxonomy ID	String	Controlled
Vaccine Year	Year for seasonal vaccines	2012		Numeric	Free text
Adjuvant	A substance added to vaccines to increase the body's immune response to the vaccine		VO	String	Controlled
Route	Eg oral, nasal spray, intradermal injection, etc.	Intramuscular, intradermal, transcutaneous	VO	String	Controlled
Scheduling	The number of times a substance is administered within a specific time period.	e.g. prime / boost scheme		String	Free text
Time Point Time Point Units		1 day		Numeric String	Fixed List
Baseline Time Event	The starting point against which other events are compared	time of vaccination (first dose)		String	Free Text
Cohort Age Min Age Max	Features that describe study cohorts	e.g. antibody responders 18 45		String Numeric Numeric	Free Text
Age Units	{days, weeks, months, years}	45 years		String	Fixed List
Publication Reference	PubMed unique identifier of an article.	30843873		Numeric	Controlled
Publication Year	The year in which the study was published	2019		Numeric	
Publication Reference URL	Link to article in PubMed	https://www.ncbi.nlm.nih.gov/pubmed/	30843873	String	URL
Comparison	The contrast used for deriving the signature	1d-0d		String	Free Text
Response Behavior	Observed change in the response agent under the comparison	Up		String	Free Text

Table 2

Signature Type		Target Pathogens	Publications	Signatures		Response Components per Signature
Gene	52	38	62	480	13,812 genes	1 to 2,036 genes
Cell type	28	26	31	185	47 cell types	1 to 9 cell types
Joint	56	44	69	665		

Table 3						
Column name	Ontology	Values				
response_component	HGNC	STAT1, IRF9, SPI1, CD74, HLA-E, TNFSF13B				
tissue_type	Cell Ontology	peripheral blood mononuclear cell				
exposure_material	Vaccine Ontology	VO:0000045; VO:0000046 (Fluarix; Fluvirin)				
target_pathogen	NCBI Taxonomy	Influenza A virus (A/Brisbane/59/2007(H1N1)); Influenza A virus (A/Brisbane/10/2007(H3N2)); Influenza B virus (B/Florida/4/2006)				
vaccine_year		2008				
time_point		1				
time_point_units		days				
baseline_time_event		time of vaccination (first dose)				
cohort		18-40 yo, subgroup high responders				
publication_reference (PMID)		21357945				
comparison		correlated with titer response index (TRI)				
response_behavior		positive				