

REVIEW ARTICLE

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Classification, Ontology, and Precision Medicine

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A GOAL OF PRECISION MEDICINE¹ IS TO STRATIFY PATIENTS IN ORDER TO improve diagnosis and medical treatment. Translational investigators are bringing to bear ever greater amounts of heterogeneous clinical data and scientific information to create classification strategies that enable the matching of intervention to underlying mechanisms of disease in subgroups of patients. Ontologies are systematic representations of knowledge that can be used to integrate and analyze large amounts of heterogeneous data, allowing precise classification of a patient. In this review, we describe ontologies and their use in computational reasoning to support precise classification of patients for diagnosis, care management, and translational research.

ABUNDANCE OF DATA

The widespread adoption of electronic health records (EHRs) affords an opportunity to collect objective and subjective observations related to demographic characteristics, findings, symptoms, diagnoses, test results, procedures, medications, nursing interventions, and so on. Very large amounts of high-throughput data, including those obtained through genomic, proteomic, and metabolomic analyses, are now being used in clinical analyses. Public data sets, such as those of the Cancer Genome Atlas and the 100,000 Genomes Project,^{2,3} provide a context (or baseline) for comparing clinical data, although such comparisons are seldom made. The volume and depth of data and the rate of its accrual are unprecedented in human history (Fig. 1).

Although EHRs document many types of data, they often impede analyses of patient-level, high-throughput or molecular data in combination with clinical data because the records are frequently incomplete, incorrect, of unknown provenance, or of insufficient level of detail. These problems are due in part to a design that is driven by billing concerns rather than a desire to document medically relevant biologic features of the patient.⁴ Data on behavioral phenotypes, environmental exposures, genome sequencing, and mobile health sensors are difficult to capture and are not systematically collected or integrated, especially since they are often “trapped” in PDF documents that can be difficult to parse into structured fields computationally.

In summary, phenotypic information about individual patients is often insufficiently detailed or inaccessible, thus obstructing the detection of similarities and the classification of patients into clinically useful groups. Such detection and classification are both challenging and important in disorders with a spectrum of symptoms, signs, biomarkers, and genotypes that may not be present in all patients, but understanding how to use these data to stratify patients and to recognize

similarities between distinct diseases is a major goal of precision medicine. For example, both *BRAF*-mutated melanoma and *BRAF*-mutated Langerhans-cell histiocytosis respond to the drug vemurafenib.⁵

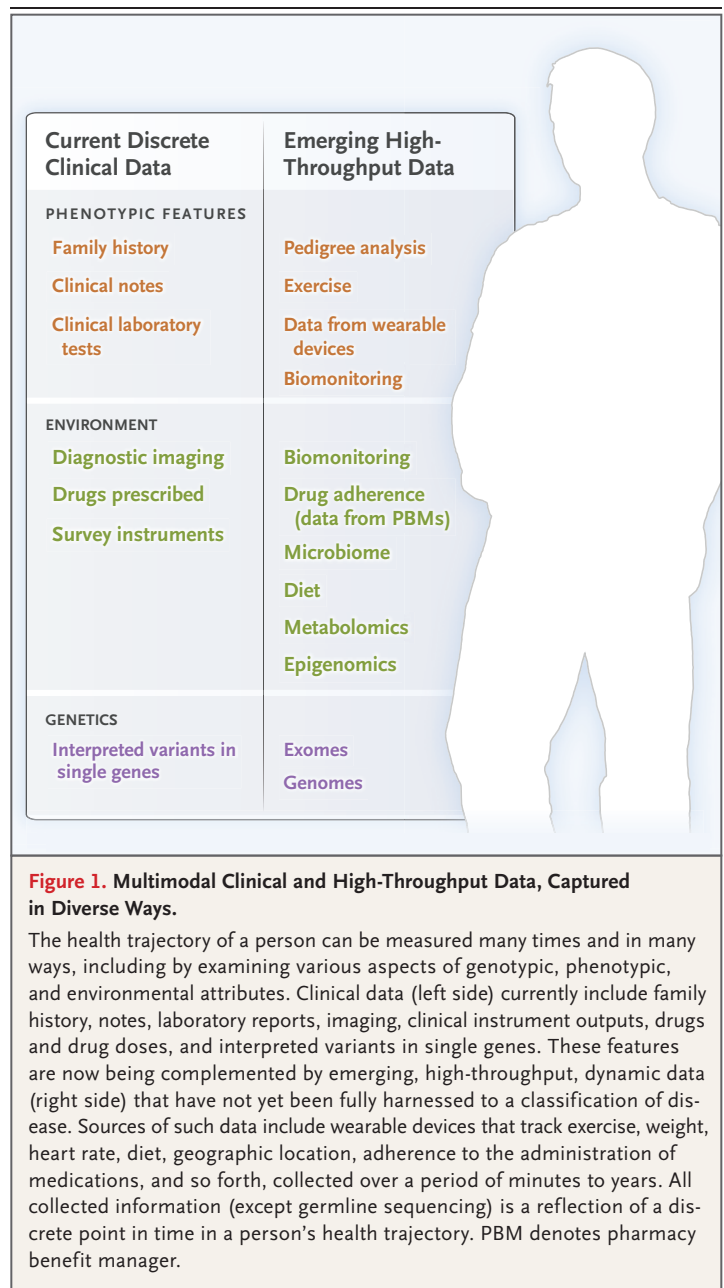
MAKING SENSE OF DATA

Data without interpretation are facts without understanding. Methods of inference, such as statistical analyses or machine learning, require categorizing subjects according to covariates, features, or both. The challenge is to create useful classifications that combine a plethora of numerical or continuous variables, dichotomies, ordinal groups, and taxonomic categories. Classifications describe entities from domains of interest, such as diseases, phenotypes, medications, and exposures, by naming the entities in each domain and providing computational specifications of differing degrees of sophistication. Increasingly formal mechanisms exist for creating such names and specifying their relationships to one another, from simple terminologies to ontologies (Table 1).

Standards exist for the majority of types of data used for clinical medicine, including diagnoses, medications, adverse reactions, procedures, laboratory data, and imaging data, as well as signs, symptoms, and other phenotypic abnormalities (Table 2). However, these data-type standards are just the first step in making data computable and patients deeply classifiable. The larger challenge is to integrate formats and structures from different sources to make them compatible.

DATA STANDARDS FOR COMPARABLE AND CONSISTENT CLASSIFICATION

Data standards can ultimately be reduced to two components: structure and semantics. Conventionally, most of us think about structure as the arrangement of data, either on an EHR screen or as a database schema behind the scenes. Semantics, in turn, refers to concepts and the relationships between them. Software systems require assertions about term equivalence. For example, without equivalent terms, clinical laboratory data with codes local to a specific laboratory or hospital are difficult to compare with corresponding data elsewhere. Similarly, diagnoses captured



in the Systematized Nomenclature of Medicine Clinical Terms (SNOMED CT, a set of terms that describe clinical findings, symptoms, diagnoses, procedures, body structures, organisms and other causes of disease, substances, pharmaceuticals, devices, and specimens)⁹ do not always correspond with similarly named conditions in the *International Classification of Diseases* (ICD).⁷ In short, mapping clinical data across systems or to the basic science data needed for mechanistic clas-

Term	Definition
Index	List of relevant terms pulled directly from a body of unstructured or semistructured text. An index is produced to improve the speed and relevance of search results.
Terminology	Set of preferred or official terms in a domain. A terminology may be a systematic nomenclature supported by a centralized body or as simple as the common usage that arises in a specific community of practice.
Thesaurus	Terminology that clusters synonyms and plesionyms (near synonyms) into categories.
Controlled vocabulary	Set of preferred terms created specifically for a domain or body of text.
Classification	Controlled vocabulary that is intended to comprehensively describe a topic or domain from a conceptual perspective and is not developed solely from a text corpus that it is meant to describe.
Statistical classification	Classification in which all concepts are mutually exclusive to avoid counting anything twice. This is typically achieved with the use of a monohierarchy, in which each concept has one and only one parent, such as the <i>International Classification of Diseases (ICD)</i> . ⁷ A statistical classification is exhaustive because it includes residual categories such as “unspecified” or “not elsewhere classified.”
Ontology	Controlled terminology invoking formal semantic relationships between and among concepts, manifested as a type of description logic, which is a subset of first-order predicate logic, chosen to accommodate computational tractability. ⁸ A common example is OWL (Web Ontology Language; www.w3.org/OWL/).

* The information presented in the table is adapted from Cornet and Chute.⁶

Type of Data	Ontology	Example of Term
Diagnoses	Systematized Nomenclature of Medicine Clinical Terms (SNOMED CT) ICD Orphanet Rare Disease Ontology (ORDO) National Cancer Institute Thesaurus (NCIT)	Triple-negative breast carcinoma (NCIT:C71732)
Phenotypic abnormalities	Human Phenotype Ontology (HPO)	Bronchopulmonary sequestration (HP:0010960)
Medications	RxNorm DrugBank ChEMBL	Panobinostat (ChEMBL483254)
Adverse reactions	Ontology of Adverse Events (OAE)	Injection-site induration (OAE:0000323)
Procedures	Medical Dictionary for Regulatory Activities (MedDRA)	Cardiac aneurysm repair (MEDDRA/10007514)
Laboratory examinations	Logical Observation Identifiers Names and Codes (LOINC)	Creatinine in serum or plasma (LOINC:2160-0)
Imaging data	Digital Imaging and Communications in Medicine (DICOM) RadLex	Periosteal cortical thinning (RID45761)

* SNOMED CT and some of the other ontologies cover multiple types of data (not shown). ChEMBL is a chemical database maintained by the European Molecular Biology Laboratory, and RadLex is a radiology-specific ontology produced by the Radiological Society of North America. Since 2015, RadLex and Logical Observation Identifiers Names and Codes (LOINC; maintained by the Regenstrief Institute) have been collaboratively producing a unified model for naming radiology procedures (<https://loinc.org/collaboration/rsna/>).

sification is often compromised by differences in naming systems or structure.

Semantics and structure are not orthogonal but deeply intertwined. One may encounter a seemingly obvious term such as “myocardial infarction” in a patient’s medical record; however, the context must be taken into account. If the mention of “myocardial infarction” was nested within an EHR partition about the patient’s family history, the interpretation would be that someone in the patient’s family, not necessarily the patient, had a myocardial infarction. Correspondingly, the clinician-reported signs and symptoms (i.e., problem list) may contain a single phrase that combines context and diagnosis — in this case, “family history of myocardial infarction.” Although this example may seem trivial, the reality is that matching of information sources must not only accommodate different semantic foundations (terminologies and classifications) but also anticipate further modification of those semantics according to their local context. Sometimes the modification is extreme, such as in the case of negation (“no history of myocardial infarction”). Exploiting the promise of precision medicine will depend on our ability to align data across patients and systems with comparable and consistent formats and contextual meaning.

FROM TERMINOLOGY TO ONTOLOGY

Terminologies have a long history of use in information retrieval (i.e., the search for documents or database entries that match certain criteria). Some of the most important resources for information retrieval in the medical domain include Medical Subject Headings (MeSH) for indexing and searching PubMed; RxNorm,¹⁰ a terminology for generic and branded drugs; and the Unified Medical Language System (UMLS), which integrates more than 100 clinical terminologies and coding systems.¹¹ These resources, which comprise standardized names and lists of synonyms and cross-references, provide the foundation for searching and indexing and are in common use in EHRs and public databases.

Ontologies differ from terminologies in that ontologies define relationships between concepts in a way that allows computational logical reasoning, enabling the drawing of conclusions

from related assertions.¹² For example, if an ontology classifies “virus” as an infectious agent and classifies “infectious meningitis” as a type of meningitis due to an infectious agent, then it would conclude that “viral meningitis” is a subclass of “infectious meningitis.” Aristotle developed conceptual taxonomies that are in some ways similar to modern bio-ontologies.¹³ More recently, scientists have used the word “ontology” to denote a computational representation describing specific domains of knowledge. An ontology consists of a set of concepts (terms) and their synonyms, as well as description-logic definitions that specify the formal relationships between the concepts (Fig. 2).

The use of description logics in an ontology can guarantee logical consistency, even with hundreds of thousands of concepts across multiple domains, enabling computational reasoning procedures to identify facts that are implied but not explicitly stated in the original data. Ontologies can thus help to leverage the latent knowledge within clinical big data by encoding the data with “computable” semantics, enabling machine-learning and other algorithms to address challenges in the analysis of multimodal, high-throughput data by integrating it with clinical meaning. Ontologies can be used in combination with natural language processing to disambiguate text concepts, such as those found in clinical notes, and improve knowledge extraction from EHRs and other sources.

Ontologies can also support integration of basic science data (e.g., data from animal models) and public knowledge (e.g., associations between genetic variants and diseases), enabling patient classification based on a corpus of data existing well beyond the EHR and permitting new clinical insights. The combination of massive data and affordable high-capacity computing provides an opportunity for unprecedented discovery of association and, increasingly, causal reasoning to gain diagnostic and therapeutic insight.

ONTOLOGIES FOR DISEASE CLASSIFICATION

The first modern medical classification that can be considered a true ontology of diseases (nosology) was developed by Carl Linnaeus (1707–1778), who divided diseases into 11 classes, 37 orders,

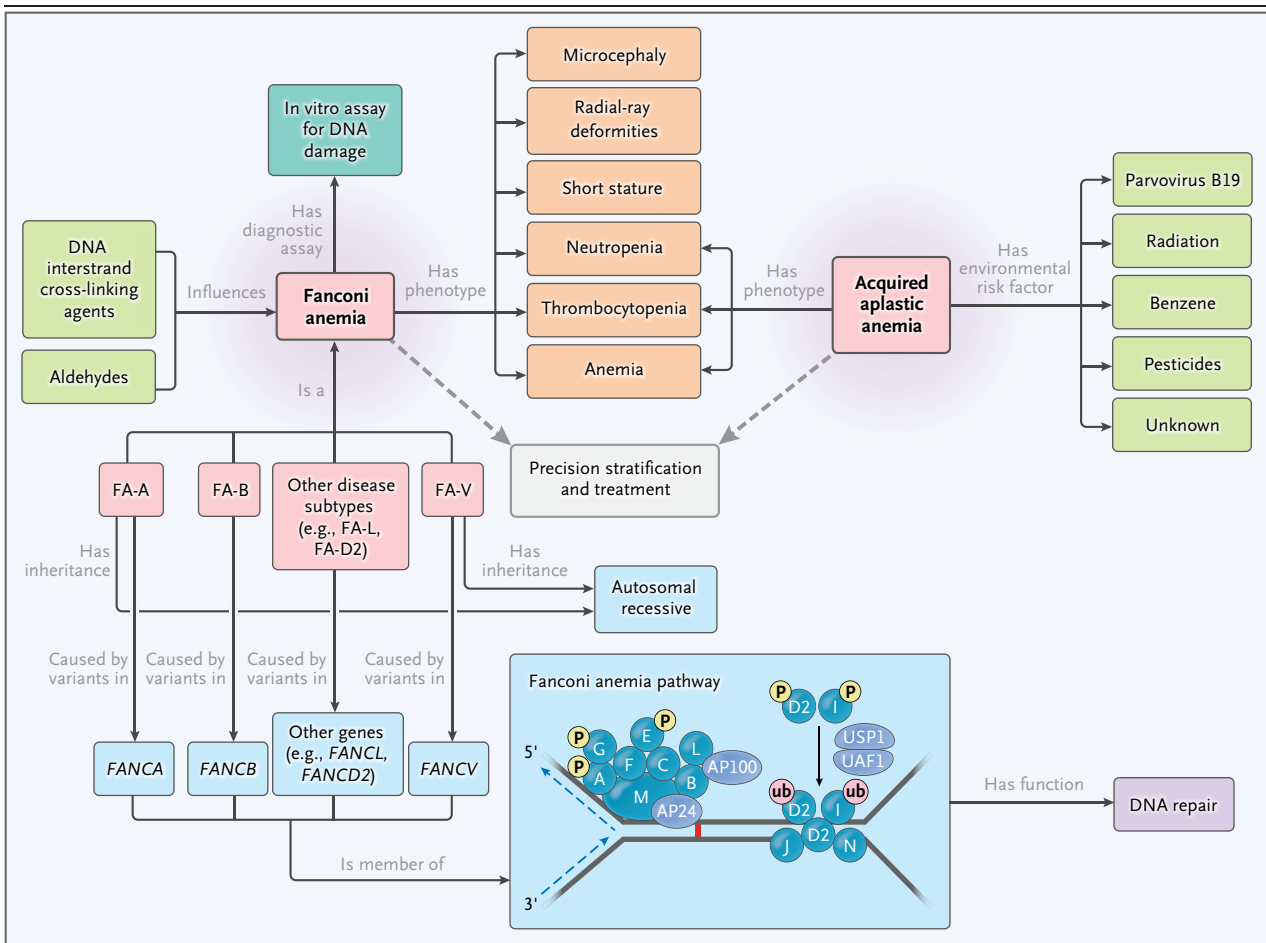


Figure 2. Ontology-Driven Representation of Fanconi Anemia and Acquired Aplastic Anemia.

Fanconi anemia and acquired aplastic anemia share several phenotypic features but have very different causal mechanisms. Computable relationships can be represented among diseases, phenotypic features, genes, and environmental exposures by interlinking terms (concepts) from sources including the Orphanet Rare Disease ontology (ORDO) (pink denotes diseases),¹⁴ the Human Phenotype Ontology (HPO)¹⁵ for phenotypic features (orange), the Chemical Entities of Biological Interest ontology for the chemical compounds (green denotes factors such as chemical exposures that can influence severity or trigger development of disease),¹⁶ and the Ontology for Biomedical Investigations for the comet assay (single-cell gel-electrophoresis assay) of DNA breakage (aqua),¹⁷ as well as the Gene Ontology (lavender denotes a biologic pathway)¹⁸ and the Reactome for molecular pathways (blue denotes disease genes and mode of inheritance).¹⁹ Ontologies can be used to support the integrative analysis of these data sources for precision stratification and treatment and to clarify underlying mechanisms, as suggested by the dashed lines. The labeled arrows between concepts represent description-logic definitions that specify the formal relationships between the concepts. FA-A denotes Fanconi anemia, complementation group A; FA-B, complementation group B; and so on. In the diagram of the Fanconi anemia pathway, the blue circles represent the proteins encoded by the Fanconi genes (e.g., A is the protein encoded by *FANCA*), and the grayish-blue ovals represent other interacting proteins; P denotes phosphorylation, and ub ubiquitination.

and 325 species. Although Linnaeus’s classification contains some errors from a modern perspective, such as the notion that leprosy can be caught by eating herring worms,²⁰ his classification laid the foundation for work that eventually led to the first edition of the ICD, in 1893.²¹ The ICD has advanced enormously since its creation as a cause-of-death inventory, but it continues to

be intended as a standard for epidemiology, health management, and billing, not as a computational representation of the patient as a biologic subject. One of the biggest issues with the ICD is that historical editions are statistical classifications that are mutually exclusive (they do not double count things) and exhaustive (they provide a place to put everything). The ICDs have

achieved exclusiveness through a monohierarchy (single parentage), with each code having one and only one parent. This precludes multiple counting but also creates arbitrary associations. For example, in the 10th edition of the ICD, malignant neoplasm of the thyroid gland (C73) is a child of malignant neoplasms but is not a child of disorders of the thyroid gland; it would be a child of both only if terms in ICD-10 could have multiple parents. Monohierarchies thus artificially constrain important axes of characterization and inquiry and impede meaningful analyses of disease and other phenotypes.

Rational disease classification dates back to Hippocrates, though it remains an active field of study today.⁶ Conventionally, phenotype has denoted observable characteristics of a person, often attributable to genotype. Increasingly, the term is being co-opted by translational researchers, working at the boundaries of “omics” data and clinical records, to define a cohort of patients with the same “diagnosis” on the basis of similar clinical and “omics” features.^{22,23}

The Electronic Medical Records and Genomics (eMERGE) study, funded by the National Human Genome Research Institute, showed the usefulness of computable phenotyping algorithms^{24,25} across medical centers with different EHR systems.²⁶ Such pragmatic and reproducible methods use standard coding systems for phenotypic abnormalities, diseases, laboratory values, medications, adverse effects, and natural language processing of free text. However, most algorithms are presented as a set of English instructions, rules, and filters, making their translation to computation and integration with multimodal, high-throughput data an exercise for each user. A limitation is that the ICD codes used in these algorithms are captured for billing purposes and not for differential diagnosis. Statistical and machine-learning approaches to cohort definitions based on mining of structured data and free-text descriptions represent another promising method of semiautomated phenotyping for large-scale case-control studies.²⁷

SNOMED CT is a compositional system, meaning that it can represent complex concepts by combining discrete facts and observations. For instance, acute perforated appendicitis can be represented by combining the concepts for “acute inflammation,” “perforation,” and “appendix structure.” SNOMED CT uses description logic

to recognize logically equivalent ways of expressing “acute perforated appendicitis,” thereby providing a consistent and computable framework.^{28,29} In this manner, SNOMED CT can be used in different ways within different systems and contexts and by different users and still result in the same conceptual and computational meaning, thus offering an advantage over simpler terminologies. Currently, however, less than a quarter of the content of SNOMED CT is logically defined; the remainder is primitive and not amenable to this method.

The increasing use of wearable health devices³⁰ and biomonitoring, as well as advances in medical digital imaging, portends growth in the volume of clinical data. Ontologies can help to organize and analyze vast quantities of data that are too large for an individual physician to manage. Patient-reported information has provided robust findings in some areas, such as genome-wide association studies.³¹ Therefore, formally encoding lay synonyms of medical terminology within an ontology³² may improve our ability to classify patients in meaningful disease groupings by integrating patient-reported information with standard medical terminology.

RARE DISEASES AND THE HUMAN PHENOTYPE ONTOLOGY

Ontologies have made a substantial contribution to translational research and the genetic diagnosis of rare disease. The sequencing of exomes and genomes has enabled the discovery of hundreds of novel disease-associated genes, and the diagnostic yield (percentage of patients who receive a molecular diagnosis) in many large-scale exome- or genome-sequencing studies is now at least 35% for some disease groups.³³⁻³⁷ In some cases, the diagnosis results in a change in clinical management, as well as family counseling.³⁸⁻⁴⁰

Variant-driven analysis aims to identify a disease-causing variant among the roughly 25,000 to 100,000 variants in a typical exome or approximately 4.5 million variants in a typical genome. Realizing the full clinical value of these data requires additional information about diseases and phenotypic abnormalities. Ontologies provide a computational-analysis framework that contextualizes the molecular data within an evaluation of the phenotypic presentation. The Human Phenotype Ontology (HPO) enables a deep pheno-

Wiedemann–Steiner Syndrome Profile	Patient 1 Profile (3-year-old girl)	Patient 2 Profile (14-year-old boy)
DIGITS		
Short toe HP:0001831	<input type="radio"/> None	<input checked="" type="radio"/> Long toe HP:0010511
Short middle phalanx of finger HP:0005819	<input checked="" type="radio"/> Cone-shaped epiphysis of the phalanges of the hand HP:0010230	<input type="radio"/> None
DEVELOPMENT		
Developmental delay in speech and language HP:0000750	<input checked="" type="radio"/> Developmental delay in speech and language HP:0000750	<input checked="" type="radio"/> Developmental delay in speech and language HP:0000750
Intellectual disability HP:0001249	<input checked="" type="radio"/> Global developmental delay HP:0001263	<input checked="" type="radio"/> Global developmental delay HP:0001263
SKELETAL		
Microcephaly HP:0000252	<input checked="" type="radio"/> Microcephaly HP:0000252	<input checked="" type="radio"/> Macrocephaly HP:0000256
Short stature HP:0004322	<input checked="" type="radio"/> Proportionate short stature HP:0003508	<input type="radio"/> None
FACIAL		
Thin upper lip HP:0000219	<input checked="" type="radio"/> Thick upper lip HP:0000215	<input type="radio"/> None
Hypertelorism HP:0000316	<input checked="" type="radio"/> Hypertelorism HP:0000316	<input type="radio"/> None
Blepharophimosis HP:0000581	<input type="radio"/> None	<input checked="" type="radio"/> Blepharophimosis HP:0000581
Epicanthus HP:0000286	<input type="radio"/> None	<input checked="" type="radio"/> Epicanthus inversus HP:0000537

Figure 3. “Fuzzy” Matching of Phenotypic Profiles.

Shown in this example are portions of the HPO profiles (green and brown) of two patients for whom clinical dysmorphologic analysis did not help establish the diagnosis, even though they were seen in the same clinic within weeks of each other. Clinical exome sequencing showed a mutation in *KMT2A* in both patients, which in combination with the phenotype comparisons led to a diagnosis of the Wiedemann–Steiner syndrome.⁴² Each set of HPO terms is compared with all other phenotypic profiles in the HPO database to find the best nonexact (“fuzzy”) match.⁴¹ Each patient has a distinct phenotypic profile that only partially matches the computational model of the Wiedemann–Steiner syndrome derived from the literature (blue). Patient 1 had microcephaly, whereas Patient 2 was found to have macrocephaly; microcephaly had been observed in only one previously described patient.⁴³ Some of the matches are relatively specific for this syndrome, such as blepharophimosis, and contribute more to the matching score than do features that are common to many diseases, such as intellectual disability. The final matching score can be calculated from the matching score for each query term and represents the proximity of the query term to its best match in the computational disease definition. A perfect match between a phenotypic feature of a patient and a feature of the disease is symbolized here by a black circle and would be assigned a high match score. A nonexact match (a fuzzy match, in which the patient has a feature that is similar to a feature of the disease but is not an exact match) is symbolized by a circle that is half black and half white and would be assigned a lower matching score. A lack of a match (a patient who does not have a feature that characterizes a disease) is symbolized by a white circle and can be penalized by some computational similarity algorithms. Existing algorithms exploit information in the ontology and annotations in many ways; commonly, they take into account the specificity of the term, usually calculated as the information content (not shown).

typing approach wherein computable phenotypic profiles of human diseases and individual patients allow the linking of terms that are close to one another in the hierarchy and weighted according to the specificity of individual phenotypic abnormalities^{15,41} (Fig. 3). There are more

than 13,500 terms in a subtype hierarchy, which also encodes the specificity of each term for differential diagnosis as a function of the frequency of the term across all diseases in the HPO database. For example, the Marfan syndrome is characterized by relatively specific HPO terms

such as ectopia lentis, which is found in 40 mendelian diseases in the HPO database, as well as less specific terms such as scoliosis, which is a characteristic of 424 mendelian diseases.

The HPO differs from other clinical terminologies and ontologies in that it provides a substantially more detailed representation of clinical phenotypes⁴⁴ and it is designed for computational analysis by linking to computational disease definitions and to ontologies of gene function, anatomy, biochemistry, and other biologic attributes.¹⁵ For instance, “neutropenia” is logically defined with the use of terms from three ontologies covering the domains of cell types, gross anatomy, and attributes: “neutrophil,” “blood,” and “decreased count.” With the use of this multi-attribute classification, a matching procedure can be applied to every disease in the database to find the closest match, or it can assist phenotype-clustering efforts by finding patients with similar phenotypic manifestations. Because this method represents the patient as a biologic subject, the Monarch Initiative has extended the approach to human–mouse phenotype comparisons on the basis of a cross-species ontology of anatomy^{45,46} and the Mammalian Phenotype Ontology.⁴⁷⁻⁴⁹

HPO-based phenotypic profile matching used in combination with genome sequencing has allowed prioritization of candidate genes with predicted pathogenic variants.⁵⁰⁻⁵⁴ This approach helped establish diagnoses in 28% of children who might have otherwise remained on “diagnostic odysseys.”⁴² Use of the HPO and data from animal models yielded about 10 to 20% more molecular diagnoses than those obtained by manual curation of sequence data from patients in the Undiagnosed Diseases Program.⁵⁵ Similarly, a machine-learning approach was applied to a cohort of 2045 persons with bleeding and platelet disorders to identify novel “fuzzy” phenotypic profiles associated with rare pathogenic variants, enabling the identification of disease-associated genes.⁵⁶⁻⁵⁸

THE FUTURE OF ONTOLOGIES IN MEDICINE

The ability to leverage extremely large amounts of data in order to answer the question, “Have I seen a case like that?” and to identify effective, safe treatments is a long-cherished aspiration.⁵⁹ But what does it mean to be the patient in “a

case like that”? Which axes of patients’ characteristics bear scrutiny: demographic data, signs, symptoms, family history, diagnoses, anthropometrics, test results, radiographic studies, or “omics” measures? How much of this information is already in my patient’s record? How much is in the records of putative patients similar to mine? How big is the universe of corresponding data that I can examine, in my practice, in my hospital, in my group, in my state, in my country? This is the logical extension of the “learning health system,”⁶⁰ taken to the level of nearly homogeneous groups of patients. The National Research Council convened a forum on precision medicine,¹ which proposed a “new taxonomy” for biology and medicine that would be structured to recognize and avail the multiple axes of basic science and clinical characteristics as a matrix defining disease endotypes (Fig. 4). The annotation of data with ontologic tags must also become an automated, background task; this is already a reality for some data sources^{61,62} and is tantalizingly close for others.

What are the barriers blocking progress toward the goal of individualized medicine? The first and most challenging is a set of privacy laws, enacted without consideration for the data-rich and data-dependent world in which we now find ourselves. Patient privacy and confidentiality are necessary to maintain the cooperation of our patients and their trust in us. Nevertheless, a new ethical framework may be in order, balancing the needs of society and future patients with legitimate expectations of privacy⁶³ and the wishes of those who want to share their medical data for the betterment of society.⁶⁴

The second barrier is the cost and effort of getting data into and out of EHRs. Manual input of structured data by clinicians is not scalable and is not a good use of clinicians’ time. Emerging efforts on standard application interfaces with EHRs from devices and data sources⁶⁵ could help, as could patient-collected and patient-entered information.^{32,66,67} Systematically harvesting signs, symptoms, severity, and other clinical details from dictated notes or even from audio capture of the patient encounter is becoming increasingly practical.⁶⁸ The third barrier is a lack of comparability and consistency among data and knowledge resources (e.g., public databases and clinical references, EHR systems and implementations, and clinical laboratories), which trans-

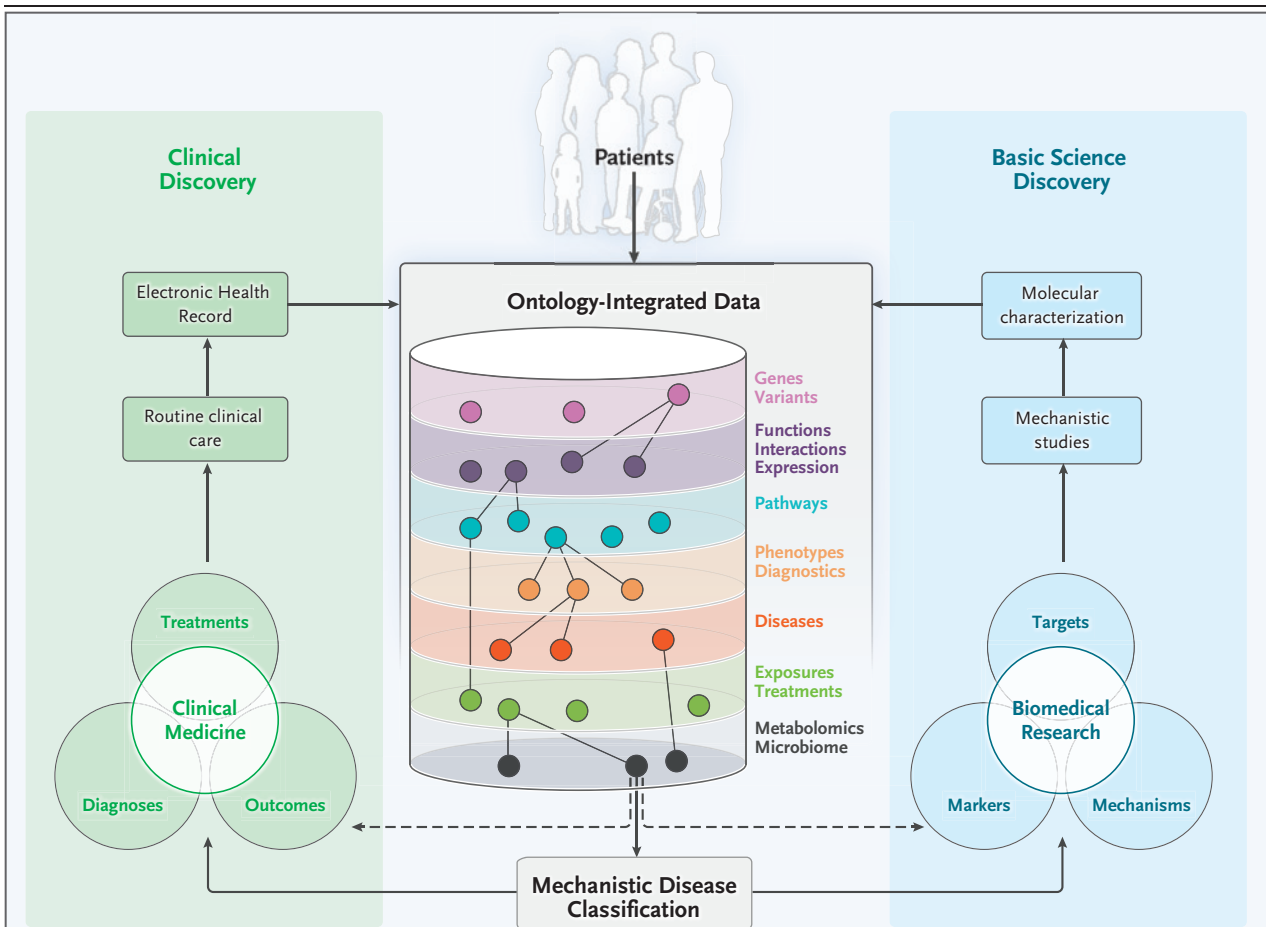


Figure 4. Ontology-Based Mechanistic Classification of Disease.

Well-structured clinical data can be readily integrated with discovery research data by using ontologies, which make clinical and basic science observations “computable” in a way that reflects present knowledge and allows new inferences. Integrating the two streams of data enables a mechanistic classification of disease across many data types, making a more refined and dynamic classification of patients possible.¹

lates to a lack of interoperability. Harmonization can best be achieved through consistent investment and community participation in computational resources for translational research.^{69,70}

Disclosure forms provided by the authors are available with the full text of this article at NEJM.org.

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