Diagnostic strategies in patients with undiagnosed and rare diseases

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Abstract

Rare diseases are life-threatening or chronically debilitating conditions affecting millions of people worldwide. In many instances, the patients experience a delay in their diagnosis or remain undiagnosed despite extensive investigations carried out by specialists. There are several explanations to account for this phenomenon including the socioeconomic context and the lack of an established consensus for diagnostic testing. Nonetheless, the widespread use of genetic and genomic tests in the past decades has had a major impact on clinical reasoning paradigms, and new troves of data are constantly being generated and analyzed. This requires constantly updating tools to match the discovery rate and allow reanalysis. In this review, we summarize the latest international recommendations and guidelines to address the problem of diagnostic deficit as well as present the current diagnostic workflows. Increasing access to exome and genome sequencing technologies and biological validation, gaining insight into the interpretation of multi-omics datasets, and fostering data sharing would reduce the long diagnostic odyssey and diagnostic gap.
INTRODUCTION

In this review, we summarize the latest international recommendations and guidelines to classify undiagnosed patients as well as present the current diagnostic workflows. We depict the advanced sequencing techniques revolutionizing genetic diagnostic practices, the future of multiple omics technologies (such as epigenomics, transcriptomics, proteomics, and metabolomics), and the use of *in silico* prediction of variant pathogenicity and functional genomics.

Rare diseases (RDs) are very numerous (more than 6000), with many of them being ultra-rare. In the European Union (EU), the definition of RDs was established in the EU Regulation on orphan medicinal products (1999) as life-threatening or chronically debilitating conditions affecting no more than 5 in 10,000 individuals[1]. The American Orphan Drug Act (1983) defined RDs as disorders affecting fewer than 200,000 individuals in the United States (US)[2]. Nguengang et al. calculated that 71.9% of RDs are genetic (including all diseases known or suspected to be familial), and 69.9% are exclusively pediatric onset[3]. Currently, the estimated prevalence of RDs is at least 3.5%-5.9%, which equates to 263-446 million individuals worldwide[3]. Moreover, RDs pose an economic burden as direct medical costs per patient are estimated to be around 3-5 times higher than controls of the same age without RDs[4].

The high frequency with which RDs remain undiagnosed[3] is a major challenge, as reflected in one of the International Rare Diseases Research Consortium (IRDiRC) goals for 2017-2027, which encourages to achieve a diagnosis of patients within one year if their disorder is known in the medical literature[6]. The term diagnostic odyssey refers to the time between when a potential RD is noted until the final diagnosis is made, while diagnostic impasse refers to the difficulty in achieving a diagnosis after performing all currently available procedures. The diagnostic deficit is usually associated with patients with complex phenotypes, the lack of genotype/phenotype correlation, or the lack of certainty of the clinical impact of a given genetic variant. Diagnosis in RD patients not only provides answers for patients and families but also has a potential clinical impact, which includes gaining knowledge on the natural history of disease and prognosis, providing genetic counseling, guiding personalized treatments, offering patient support networks, enabling participation in research studies, informing reproductive choices, and impacting the health of relatives[7].

Many but not all patients with an undiagnosed disease have an RD[8]. Undiagnosed and rare diseases (URDs) are conditions that elude diagnosis by a referring specialist or center of expertise for a long time and despite extensive state-of-the-art investigations[8]. Graessner et al. estimated that around 50% of patients with RDs remain undiagnosed even in advanced expert clinical settings where genome sequencing techniques are applied routinely[9]. Available investigations vary in each socioeconomic context, and there is not a consensus list of laboratory and ancillary tests to be performed before concluding that a patient is undiagnosed. Our ability to diagnose URDs is limited by our incomplete knowledge of the natural history and clinical expression of the disease, the genotype/phenotype correlations, the full mutational spectrum associated with all RDs, and the number of unique RDs that have yet to be discovered[10]. The IRDiRC Solving the Unsolved Task Force proposes to classify URD patients into specific subsets with significant utility for optimizing diagnostic strategies [Table 1][10]. However, in many cases, it is more important to take into account the potential explanations for the diagnostic deficit before undergoing follow-up tests. In these cases, we can use an alternative classification system [Table 2].
Table 1. Clinical groupings and diagnostic strategies for patients with undiagnosed and rare diseases (adapted from the IRDiRC solving the Unsolved Task Force\cite{10})

| Clinical groupings                                      | Recommended diagnostic strategies                                                                                                                                 |
|--------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Patients with clinically recognizable disorders         | No causative variant after an appropriate highly sensitive test (e.g., single-gene disorders such as neurofibromatosis type 1)                                      |
|                                                        | Explore new tests and computational tools (e.g., RNAseq might be useful in patients with suspected neurofibromatosis type 1 and negative conventional tests)      |
|                                                        | No identified causative variant in the context of genetic heterogeneity (e.g., retinitis pigmentosa)                                                                 |
|                                                        | Get large datasets of patients, detailed phenotypic and genomic information, and share data (data sharing of patients with a similar phenotype can help in the identification of disease-causing variants in yet-to-be-discovered disease genes) |
|                                                        | Explore new tests and computational tools                                                                                                                                 |
| Unsolved but recognizable disorder (e.g., VACTERL association) |                                                                                                                                                             |
| Patients without clinically recognizable disorders      | Patients with syndromes without a name (SWAN), which are not recognizable as a previously described disorder                                                     |
|                                                        | Exome or genome sequencing as first-line tests                                                                                                                                                                    |

The widespread use of genetic tests in the clinical setting has had a great impact on the paradigms of clinical reasoning in the field of clinical genetics. Classical linear reasoning, in which phenotype assessment leads to a clinical suspicion that is further confirmed or refuted by genetic testing, has increasingly been replaced by circular reasoning, where both phenotype and genotype are assessed in parallel\cite{14,15}. Moreover, some diagnoses are only considered after the identification of a specific genetic variant. However, considering phenotype assessment and clinical suspicion is still an essential piece of the diagnostic process.

**THE GENOMIC SCENARIO: EXOME AND GENOME SEQUENCING**

Exome and genome sequencing (ES and GS, respectively) are now recommended by the American College of Medical Genetics and Genomics (ACMG) to be considered as first- or second-tier tests (after chromosomal microarray (CMA) or focused gene testing) for patients with congenital anomalies (CA), developmental delay (DD), or intellectual disability (ID)\cite{16}. Only patients with clinical presentations suggestive of a specific diagnosis should undergo targeted testing first\cite{16}. This may include patients with suspicion of a chromosomal disorder, a diagnosis in which sequencing may not be diagnostic (e.g., fragile X syndrome), or known family history of a disorder\cite{16}. This recommendation supersedes the previous advice to perform CMA as first-line tests in patients with CA and DD/ID (setting aside patients with autism spectrum disorder without other delays)\cite{17}.

The recent change in recommendations is due to the high diagnostic yield of ES and GS. The Ontario Health Technology Assessment reported a diagnostic yield of 43% for GS and 34% for ES (not differentiating between the trio and singleton testing), compared with the diagnostic yield of 21% for standard genetic testing (which typically included CMA, candidate single-gene testing, or large gene panel testing)\cite{18}. Although GS, by definition, captures more comprehensively all types of variants in testing, ES is expected to be used for a long time in clinical genetics given the lower cost, focused approach, and reduced burden on downstream analysis compared to GS\cite{19}. When performing ES and GS, best practice includes trio testing if available to help contextualize rare variants, but they also can be effectively performed as singleton testing, with diagnostic yield being slightly reduced\cite{16}.

Although first-line ES or GS has still not been established as standard clinical practice in all settings, the increased use of ES in the past decade has had a significant impact in the field of URDs. If a patient remains undiagnosed after performing ES, it is essential to know the limitations of ES to continue with the diagnostic process [Table 3]. In addition to recommending ES/GS as first-tier tests (or second-tier tests in certain cases), the ACMG suggests a diagnostic algorithm for the evaluation of patients with GDD/ID if ES
Table 2. Potential explanations of the diagnostic deficit and diagnostic strategies

| Potential explanations                                                                 | Recommended diagnostic strategies                                                                 |
|----------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|
| No variant has been identified                                                         | • Precision phenotype assessment; includes deep phenotyping using Human Phenotype Ontology (HPO) terminology [11] |
| • The disorder has not been recognized by the referring specialist (because of recent discovery, atypical presentation, concomitant comorbidities, and/or lack of expertise), and thus the appropriate analysis of existing data (e.g., focused on candidate genes) or the appropriate test (e.g., Angelman syndrome methylation analysis) has not been performed | • Literature review                                                                                     |
| • Changes in the classification of variants over time                                     | • Periodic reanalysis                                                                                  |
| • Limitations of current testing methodologies [Table 3]                                 | • Consider alternative tests/new computational tools and tests                                        |
| • Difficulties in the interpretation of complex inheritance patterns (e.g., genetic modifiers and polygenic inheritance) | • No general recommendation can be made. Genome-wide association studies can be used to identify single nucleotide polymorphisms significantly associated with a complex trait (e.g., human height)[12] |

A variant has been identified (adapted from Pijuan et al. 2021)[13]

A variant has been identified in one of the known disease genes:
• The genotype matches the phenotype (at least partially), but there is insufficient evidence for candidate variant (variant of unknown significance)
• The genotype does not match the phenotype:
  - Difficulties in the interpretation of reduced penetrance and variable expressivity of known disorders
  - Yet-to-be discovered allelic disease*
• A variant has been identified in a yet-to-be-discovered disease gene, but there is insufficient evidence for candidate variant or gene causality*

*In any group of patients with a clinically recognizable disorder or suspected new disease, consider a large dataset of patients, detailed phenotypic and genomic information, and data sharing.

is nondiagnostic[20]. The next step would be CMA, and, in the case it is nondiagnostic, then the clinician could consider further evaluation/testing including periodic ES reanalysis every 1-3 years, fragile X syndrome testing, metabolic testing and/or mitochondrial DNA sequencing depending on clinical presentation (although mtDNA analysis is sometimes performed with ES), and karyotyping to assess for abnormal segregation of balanced chromosomal rearrangements[20].

CMA is still the standard of care for detecting copy number variants (CNVs) in clinical laboratories[17]. However, many efforts have been made to develop ES-based CNV detection tools, such as the modified ExomeDepth workflow[19]. It is expected that GS will eventually replace CMA as the gold standard for CNV detection, considering that it covers both coding and noncoding regions. The next frontier in next-generation sequencing (NGS) technologies is long-read GS, which might outperform the limitations of short-read sequencing concerning the identification of structural variants, sequencing repetitive regions, phasing of alleles, and distinguishing highly homologous genomic regions[23]. The main limitations of current long-read NGS technologies are difficulties in library preparation, higher error rate compared to short-read NGS, higher costs, and difficulties in data analysis and storage[23].
Table 3. Types of variants missed by ES and recommended diagnostic strategies[^10-22]

| Types of variants missed | Recommended diagnostic strategies |
|--------------------------|----------------------------------|
| • Large CNVs (except when specifically included in analysis pipeline) | CMA |
|                          | GS |
| • Small CNVs (except when specifically included in analysis pipeline) | MLPA (targeted approach) |
|                          | High-resolution CMA |
|                          | GS |
| • Balanced chromosomal rearrangements | Karyotype |
|                          | GS |
| • Low coverage regions | Deep sequencing of multiple tissues |
| • Low-level mosaicism | Repeat expansion testing (targeted approach) |
|                          | Long-read GS |
| • Repeat expansions | Sanger sequencing of small fragments of genome (targeted approach) |
|                          | GS |
|                          | RNA-seq |
| • Splicing mutations (synonymous, splice site, or intronic mutations) | GS |
|                          | RNA-seq |
| • Regulatory DNA mutations (promoter, enhancer, and others) | Methylation arrays |
| • Imprinting | New computational tools |
| • Transposable elements (retrotransposons) |

CMA: Chromosomal microarray; CNVs: copy number variants; GS: genome sequencing; MLPA: multiplex ligation-dependent probe amplification; RNA-seq: RNA sequencing.

Genomic data reanalysis

The adoption of ES has accelerated the rate of novel gene discovery for Mendelian conditions. The annual number of discoveries of genes underlying RDs peaked between 2012 and 2015 (approximately 285 per year), and it has declined slightly thereafter[^24]. This is one of the arguments in favor of periodic reanalysis of ES/GS data, but it is only the tip of the iceberg. There are several other reasons causative mutations might be unrecognized. Data are usually analyzed according to the reported patient phenotype, and key elements might not be available to the clinical laboratory or might not have emerged at the time of the first analysis[^25]. The constantly growing knowledge of gene networks is a useful resource for hypothesis generation and prioritizing candidate genes. The processes of both calling variants from short sequence reads and to annotating the impact of variants are performed with imperfect and constantly evolving bioinformatics tools[^25,26]. Similarly, current phenotype-driven genomic diagnostics software (which usually uses HPO terminology) and databases that collect published mutations (e.g., Human Gene Mutation Database®) are incomplete[^25]. Besides, a significant amount of time is required for an expert geneticist to analyze ES data, and there are various sources of variability and bias that make it difficult to exactly replicate the analysis[^25]. According to various reports, reanalysis of ES data may enhance the diagnostic yield by 10%-18.9%^[^25,27-29].

Deep genomic sequencing by next-generation sequencing

It is worth noting that sensitivity for detection of mosaicism can be lower on ES/GS compared with panel testing, due to greater read depth and sequence coverage[^26]. When somatic mosaicism is suspected, the best diagnostic strategy is to combine deep NGS (generally using a customized panel of candidate genes associated with the phenotype) and variant search in multiple tissues[^26]. The utility of this approach may be limited by difficulties in obtaining tissues other than blood, saliva or buccal mucosa, and skin fibroblasts for genetic analysis. Somatic mutations have been described in several noncancerous disorders, such as PI3K/AKT/mTOR pathway mutations in patients with hemimegalencephaly, GNAQ mutations in patients with Sturge-Weber syndrome, GNAS mutations in patients with McCune Albright syndrome, and NIPBL mutations in patients with Cornelia de Lange syndrome (approximately 30% of clinically diagnosed patients have somatic NIPBL mutations)^[^30,31]. Deep NGS has also been found to be useful for detecting somatic...
mutations in patients with other brain malformations, such as double-cortex syndrome or polymicrogyria\cite{32,33}.

**Classical karyotype studies**

Despite new technological advances, it should be noted that chromosome analysis still has a role in the study of patients with URDs. Chromosome analysis remains particularly valuable in the detection of mosaic abnormalities; the clarification of unbalanced translocations, rings, and complex rearrangements; and the detection of balanced rearrangements\cite{34}. Patients with *de novo* apparently balanced rearrangements might harbor cryptic deletions or potential gene disruptions, which would require further testing\cite{35}.

**MULTI-OMICS APPROACHES**

Massively parallel technologies appear as powerful approaches to achieve diagnosis integrated with both DNA sequencing and phenotypic data. Different biological molecules can be approached with omics techniques beyond DNA, such as RNA, metabolites, and proteins or epigenetic modifications. The analysis of these molecules offers complementary data that could help to understand diseases mechanism and provide new diagnostic tools. A general drawback of multi-omics approaches is the difficulty in obtaining the most informative tissue for the analysis when it is not blood or skin fibroblasts.

**Epigenomics**

Epigenomics is the analysis of the entire set of epigenetic modifications on the genetic material of a cell, generally intending to identify alterations in DNA methylation or histone modification in diagnosing specific disorders. DNA methylation is one of the most commonly studied epigenetic modifications, and aberrant DNA methylation has long been associated with several RDs, such as Angelman and fragile X syndromes. Several methods have been developed to assess genome-wide DNA methylation in peripheral blood\cite{36-38}. These methods have proven particularly useful to diagnose patients for whom there is a clinical suspicion of an imprinting disorder or Mendelian disorders caused by mutations in genes that code for proteins involved in epigenetics (e.g., Coffin-Siris, Kabuki, and Sotos syndromes)\cite{36,37}. The potential utility of diagnostic clinical genomic DNA methylation testing in patients with RDs has been recently reported\cite{38}. Using a clinical genome-wide methylation test for patients with RDs, 35% of patients in a targeted cohort were positive for a diagnostic episignature versus 11% of patients in a screening cohort, proving that this approach can be applied for the untargeted study of patients with URDs\cite{38}.

**Transcriptomics**

The utility of analyzing the patient’s transcriptome for RDs has already been proven. Single-gene RNA sequencing has been carried out to elucidate the role of candidate genetic variants of uncertain significance (VUS) by combining reverse transcriptase to create cDNA and RT-PCR to measure RNA\cite{39}. Performing RNA sequencing (RNA-seq) of the entire transcriptome in a single run can facilitate the detection of aberrant expression, aberrant splicing, and mono-allelic expression\cite{40}. RNA-seq has proven particularly useful in specific clinical settings, such as the analysis of muscle biopsies from patients with rare muscle disorders and cultured fibroblasts from patients with mitochondrial disorders, or recessive conditions for which only one mutation has been identified\cite{41-43}. Some studies have assessed the utility of RNA-seq, in combination with ES/GS, as a diagnostic tool for URDs of diverse disease categories, with an additional yield of 17%-18%\cite{44,45}. The main challenges of RNA-seq are the accessibility of appropriate tissues for mRNA extraction, the large amount of sequencing required to detect changes, and the necessity of large control cohorts\cite{46}. 
Proteomics and metabolomics
The analysis of proteins and the proteome can provide information about the activity, interaction, location, and composition of protein complexes of clinical interest. On the other hand, metabolomic profiles have been used as biomarkers for disease progression and response to treatment, and they are now applied in diagnostics to determine the functional consequences of a given VUS. Mass spectrometry is the most widely used platform in proteomics and metabolomics. Proteomics is the quantitative and qualitative analysis of the entire set of proteins in a given specimen, generally to identify proteins that are consistently modified or present at abnormal concentrations in specific disorders\(^{[47]}\). Proteomic strategies have been applied to investigate the pathophysiology of metabolic disorders, such as methylmalonic acidemia\(^{[48,49]}\). Metabolomics is the quantitative and qualitative analysis of all metabolites derived from sugars, lipids, proteins, and nucleic acids in a given specimen\(^{[50]}\). The most successful application of targeted metabolomics analysis is the newborn screening of inborn errors of metabolism\(^{[51]}\). Overall, significant work needs to be done to implement proteomics or metabolomics in the untargeted study of patients with URDs in routine clinical practice.

**IN SILICO BIOLOGY AND EXPERIMENTAL FUNCTIONAL STUDIES**

One of the main challenges for genetic diagnosis using NGS is the interpretation of the pathogenicity of variants, particularly when the phenotype is a source of uncertainty (e.g., reduced penetrance and variable expressivity) or when a variant is classified as VUS or localizes in a yet-to-be-discovered disease gene. Indeed, during the analysis and interpretation of NGS data in a particular patient, the probability of detecting a VUS is higher than the probability of detecting a pathogenic variant\(^{[52]}\). The ACMG has elaborated standards and guidelines to classify genetic variants into five criteria-based categories using different types of variant evidence (e.g., population data, computational data, functional data, and segregation data)\(^{[53]}\). In addition, certain bioinformatic tools facilitate the classification process (e.g., Available from: Varsome, https://varsome.com/; Franklin, Available from: https://franklin.genoox.com/clinical-db; CADD, Available from: https://cadd.gs.washington.edu/), but each geneticist must provide specific knowledge to be able to classify a certain variant\(^{[13]}\). Sometimes, segregation studies can clear up doubts, but those variants that are not reclassified remain as VUS until further studies allow their pathogenicity to be determined.

*In silico* prediction of the pathogenicity of a VUS can be improved with several tools such as literature review and data mining, pathogenicity predictors, and 3D protein modeling\(^{[13]}\). After that, functional genomics can be used for the validation of the genetic variant through molecular and cellular experiments (e.g., subcellular localization studies, expression levels, and specific studies related to protein function)\(^{[13,34]}\). At Sant Joan de Déu (SJD) Hospital and Research Institute, we developed the in-house Translational Diagnostics Program (TDP) to functionally validate both candidate VUS and variants found in patients with phenotype-genotype incongruity\(^{[13,30]}\). The TDP uses different tools of experimental and computational biology to analyze VUS and determine the function and possible pathophysiological alteration of the encoded protein. The objective is to delineate the impact of VUS by combining four stages including the in-depth and precision phenotyping with functional genomics, for the validation of the genetic variant through molecular and cellular experiments. Using this pipeline in RDs patients, we can assist the process of reclassifying variants concerning a patient’s phenotype and improve the diagnostic deficit.

**OTHER STRATEGIES FOR DIAGNOSIS AND DISEASE-RELATED GENE DISCOVERY**

In recent years, the greater integration of genomic science and medicine has made it possible to explore other strategies to achieve the diagnosis of patients or to define new genes as responsible for the disease. The director’s board of the ACMG released a position statement on how responsible sharing of genomic variant
and phenotype data is crucial to improving genetic healthcare\cite{56}. Data sharing is necessary to describe the key features of the phenotype of those with RDs, establish the association between genetic RDs and the causative genes, classify genomic variants, and improve standards used in variant classification\cite{56}. Data sharing is compatible with the imperative of protecting privacy in healthcare\cite{56}. Currently, there are online data-sharing resources such as Matchmaker Exchange, open collaboration between different platforms (including GeneMatcher, PhenomeCentral, DECIPHER, and others) to facilitate the matching of cases with similar phenotypic and genotypic profiles (matchmaking) through standardized application programming interfaces and procedural conventions\cite{57}. With the advent of multi-omics approaches, there is an increasing need to share data on tools and methods for data integration and interpretation. Regarding the discovery of new genes associated with disease, the analysis of genomic population data allows the evaluation of the strength of natural selection to identify genes and genomic regions that are constrained for variation compared to the expected mutation rates. This information reveals which genes are most intolerant to loss-of-function or missense variants\cite{58}. The predicted loss-of-function intolerance (pLI) score and the lower observed/expected upper-bound fraction (LOEUF) score can be used to identify candidate haploinsufficient disease genes\cite{57,58,59}. Another alternative strategy for the discovery of new genes is the analysis of the phenotypic effects of gene disruption using model organisms when the gene of interest is evolutionarily conserved\cite{7}. Four related projects that use animal models are the Monarch Initiative, the Mouse Genome Database, the Knockout Mouse Project, and the International Mouse Phenotyping Consortium\cite{7}. However, it should be noted that animal models often fail to recapitulate human disease phenotypes. Other options are the possibility of deriving human-induced pluripotent stem cells (hiPSCs) from patient cells (e.g., fibroblasts) or generating pathogenic variants in wild-type hiPSCs using editing approaches (e.g., CRISPR/Cas9 technologies), which could be coupled with the generation of 3D organoids.

**CONCLUSION**

We are currently witnessing how genetics and genomics of URDs are one of the fields in which precision medicine and translational research are opening new paths and opportunities in etiological diagnosis. Advances in genetic and genomic testing have markedly improved the rate and time to diagnosis of patients with URDs. However, it is important to note that all techniques have limitations. Novel multi-omics techniques are rapidly advancing toward clinical practice, and in silico studies and functional analyses allow us to validate the significance of the findings.

Increasing access to exome and genome sequencing technologies and biological validation, gaining insight into the interpretation of multi-omics datasets, and fostering data sharing would reduce the long diagnostic odyssey and diagnostic gap.

**DECLARATIONS**

**Authors’ contributions**

Conception or design of the work, drafting the article, critical revision of the article, final approval of the version to be published: Casas-Alba D, Pijuan J

Conception or design of the work, critical revision of the article, final approval of the version to be published: Hoenicka J, Vilanova-Adell A, Vega-Hanna L, Palau F

**Availability of data and materials**

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Conflicts of interest
All authors declared that there are no conflicts of interest.

Ethical approval and consent to participate
Not applicable.

Consent for publication
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