Implementing ACMG guidelines on sequence variant interpretation: software-assisted variant curation and filtering

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Introduction

Since their publication in 2015, the American College of Medical Genetics and Genomics' standards and guidelines have been widely adopted. The College made these available as an educational resource for clinical laboratory geneticists to help them in the interpretation of sequence variants. Although adherence to these standards and guidelines is voluntary and cannot replace the clinical laboratory geneticist’s professional judgment, the recommendations represent a broad consensus of the clinical genetics community. With increasing volumes and the use of large gene panels (clinical, full exomes and even full genomes) in clinical genetics routine practice, labs need strong informatics tools that support them in the automation and standardization of variant assessment and reporting, in order to benefit from community standards and to keep up with the best standard of care.

“We translated the ACMG guidelines into an automated variant filtration pipeline that integrates nicely in our routine setting. When reviewing and curating variants, the software does all the time-consuming work of gathering and collating the evidence. We move straight on to reviewing the evidence and proposed classification, and confirm or modify the assessment. Our ability to implement these guidelines efficiently makes us confident we are providing the best possible quality to our referring physicians.”

Dr. Berivan Baskin (Uppsala University Hospital)

At a glance

In this white paper, you will learn:

• How Uppsala University Hospital’s Molecular Genetics lab has implemented the ACMG guidelines and standards on sequence variant interpretation.

• How the Cartagenia Bench NGS platform facilitates automation of variant classification pipelines.

• How the ACMG classification functionality enables evaluation of all ACMG criteria on a single variant level providing a complete evidence overview.
In this whitepaper, we showcase how Cartagenia Bench Lab NGS enables labs to implement their take on the ACMG recommendations. The Molecular Genetics department at Uppsala University Hospital illustrates how it is has implemented the recommendations in their specific routine diagnostic setting, using a flexible, drag-and-drop interface to build and store the lab’s variant triage protocol.

Key requirements

The standards and guidelines describe an evidence-based approach for the assessment of variants of clinically validated genes. The recommendations use literature and database-based criteria to classify variants in five different categories: benign, likely benign, uncertain significance, likely pathogenic and pathogenic. Evidence levels are weighted (e.g. “Strong”, “Moderate”). To allow labs to automate their implementation of this evidence-based approach with the possibility to manually review the results, a number of specific tools are required.

Based on the guidelines we have grouped the 28 criteria into 11 categories where for each category the corresponding evidence is given to optimally support the variant assessment process. Depending on the lab specialists’ acceptance of criteria, a classification will be predicted according to the scoring matrix as proposed in Richards et al., 2015. The possibility to take along the variant classifications and summaries based on the ACMG criteria, combined with Bench Lab’s flexible analysis approach provides a time-efficient way to address variant management and curation challenges.

Annotation sources, such as population, disease-specific, and sequence databases

The guidelines recommend the use of a wide range of criteria. Examples include: population databases such as the Exome Aggregation Consortium (ExAC, http://exac.broadinstitute.org/), Disease databases such as ClinVar (http://www.ncbi.nlm.nih.gov/clinvar), and sequence databases such as RefSeq (http://www.ncbi.nlm.nih.gov/refseq/rsg). With Cartagenia’s Bench NGS platform, labs can integrate and use a wide range of community-accepted resources, including the tools and data sources recommended in the ACMG guidelines. Moreover, with Bench NGS, labs can benefit from full version control and traceability on these resources.

Rules for combining criteria to classify sequence variants: automation of decision trees and scoring combined with variant review to manually accept ACMG criteria

The guidelines recommend a broad set of informative criteria for assessing the clinical impact of a sequence variant. With each criterion, they also provide a level of evidence strength. For example, for a de novo variant in a patient with the disease, no family history and with both maternity and paternity confirmed, the evidence to classify the variant as ‘Pathogenic’ is suggested to be ‘Strong’. Other levels are ‘Very Strong’, ‘Moderate’ and ‘Supporting’. The guidelines also propose a scheme of rules by which labs can combine different criteria for classifying variants with different levels of evidence. In order to automate such a scheme, a tool set is required to represent rules into a workflow, and associate scores to variants accordingly. Bench NGS elegantly provides such a system by means of classification trees. The user can choose from a library of filter components that each represent filter criteria such as ‘population frequency’, and can drag-and-drop these into a decision tree.
Review PROPOSED CLASSIFICATION
+ Include variant in REPORT
+ Add to MANAGED VARIANT LIST (MVL)

Triage and classify variants with CLASSIFICATION TREES

MANUALLY ACCEPT ACMG CRITERIA for each variant based on available evidence
After the classification tree has been executed, variants marked for review can be manually curated. This means the ACMG criteria can be evaluated to be accepted or rejected for each variant and based on the accumulated evidence a classification (e.g., likely pathogenic) can be assigned to this particular variant.

In figure 3, an overview of the ACMG classification criteria is given for the p.Trp1105Ser variant. At a glance, we can see the labels that were associated to this variant by the decision tree while within the ACMG Guidelines tab, we can get insight into the different evaluation criteria accompanied by the relevant evidence summaries. When evaluating for example the computational evidence, we can decide to accept the PP3 criterium for this variant by taking into account the outcomes from multiple functional effect prediction algorithms, conservation and evolutionary splicing impact. At the bottom of figure 3, we can immediately see the majority of outcomes is shown in red and therefore the overall prediction is (probably) damaging. Besides the color coding the score ranges can also be visualized on hover and together they enable a quick and easy evaluation to accept the PP3 criterium for computational evidence.

When completing the evidence assessment for the ACMG guidelines a variant classification will be proposed.

In figure 3, after accepting the PP3 criterium and rejecting BP4, a temporary VOUS classification is proposed. If this classification seems justified, it can be copied to the variant’s classification right away and if desired a summary can be copied to the variant’s assessment. Moreover, if this variant would have been observed previously and an ACMG scoring had been performed, its classification can easily be retrieved, without the need to go through the same process again.

**Figure 2.** An example decision tree. This tree shows how a number of variants are checked against common population databases, and - based on a specific Minor Allele Frequency - are labeled as “benign” with ACMG category “BA1”, as per the recommendation to assert an allele frequency over 5% in the Exome Sequencing Project, the 1000 Genomes Project, or the Exome Aggregation Consortium. The path a selected variant follows through the tree is highlighted. As a variant is filtered through the tree, it picks up annotations, classifications, scores and labels as configured by the lab.
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Figure 3. The ACMG variant classification tab provides insight into the different evaluation criteria accompanied by the relevant evidence summaries. Based on the provided evidence, ACMG criteria can be accepted or rejected. When completing the evidence assessment for the ACMG guidelines a variant classification will be proposed.
**Implementation**

The molecular genetics laboratory at the Uppsala University Hospital has implemented the ACMG guidelines on the Cartagenia Bench NGS platform and validated their approach on a set of clinical cases. The lab has implemented different criteria as well as levels of evidence in a decision tree, partially shown in Figure 4. In this view, a validated pipeline is run on a Connective Tissue Panel sample, showcasing a variant in the COL1A2 gene that is reported as clinically relevant. The protocol represented by the tree has checked all variants in the assay, and highlighted the p.Gly949Ser variant for review. The clinical geneticist consecutively verifies relevant sources — in this case: ESP, 1000 Genomes, ExAC, HGMD, in silico score annotations from ACMG-recommended SIFT, Mutation Taster and PolyPhen, and a confirmed spectrum of missense mutations in the gene at hand. Parental samples tested negative for this variant.

**Conclusion**

With this case study, the lab has illustrated how various features of the Cartagenia Bench Lab NGS platform were used to implement an automated Standard Operating Procedure that reflects how the lab performs variant filtration. This case illustrates strong advantages in lab efficiency - whereas a manual process of variant filtration is time consuming and error prone, the lab benefits from automation of these manual protocols, freeing up time for genetic specialists to focus on variant interpretation and reporting while being able to manually review flagged variants and their associated ACMG criteria with supporting evidence. The possibility to copy the ACMG assessment from previous observations further endorses the lab specialist to avoid double work and focus on new observations. To conclude, this approach provides an elegant solution for labs to save time by automating their protocols while keeping control over the decision-making process via a dedicated variant review section.

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1. Standards and guidelines for the interpretation of sequence variants: a joint consensus recommendation of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology. Richards et al., Genetics in Medicine, advance online publication 5 March 2015. doi:10.1038/gim.2015.30

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**Figure 4.** Partial view of the Uppsala University Hospital decision tree representing their filtration strategy, investigating public and in-house variant databases, modes of inheritance, population frequency statistics databases, and variant coding effect. Top: decision tree. Middle: currently selected ACMG category PPS. Bottom: variants matching selected criteria. (Courtesy of Dr. Berivan Baskin)