Recommendations/Requirements for Molecular Proficiency Testing

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Legend of Terms

- EBV = Epstein-Barr virus
- FISH = Fluorescence in situ hybridization
- FFPE = Formalin-fixed, paraffin-embedded
- GIST = Gastrointestinal stromal tumor
- H&E = Hematoxylin and eosin stain
- HPV = Human papillomavirus
- ISH = In situ hybridization
- NGS = Next-generation sequencing
- PET = Paraffin-embedded tissue
- PT = Proficiency testing
- SNV = Single nucleotide variant
Additional Information Regarding CAP Survey Programs

• For additional information regarding the PT programs mentioned throughout these flow charts, please refer to the Surveys Catalog by clicking on the Catalog and Ordering Information link under the Laboratory Improvement header at www.cap.org.
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PT Requirements for Laboratories Accredited by the CAP

- Participation in PT is integral to the CAP’s accreditation program and is required for most tests for which the laboratory reports results.
- For analytes that require PT, each laboratory must enroll and participate in a CAP-accepted PT program. In the following flow charts, required programs/analytes will be indicated by an asterisk (*).
- For tests that do not require enrollment in a CAP-accepted PT program, the laboratory must perform an alternative assessment semiannually to determine the reliability of testing. The most common way to do this is by purchasing an external PT product, if available. Other acceptable alternative assessment procedures are split sample analysis with reference or other laboratories, split samples with an established in-house method, assayed materials or other suitable and documented means. It is the responsibility of the director to define such alternative assessment procedures and the criteria for successful performance. Any program without an asterisk (*) in the following flow charts is not a required PT program and may be used to satisfy alternative assessment requirements.
- For a full list of required programs/analyte(s), please refer to the Analyte/Procedure Index in the Surveys Catalog.
PT Referral

- The NGSST, NGSHM, and NGS-Germline programs can be used to fulfill alternative assessment requirements for laboratories performing both wet bench and bioinformatic components of the assay. If a distributive testing model is used (e.g., different parts of the NGS assay are performed by laboratories with different CLIA numbers), laboratories **cannot** use these programs for alternative assessment. To do so, laboratories would be subject to sanctions for PT referral.

- Laboratories using any other distributive testing approach must use alternative approaches to fulfill the requirement for alternative assessment. Please note that distributive testing laboratories can use PT materials for part of their laboratory quality management program; laboratories should contact CAP for additional details.
Germline Molecular Flow Chart

Are your lab activities gene-specific (eg, cystic fibrosis CFTR screening)?

Yes

Is the analyte(s) required by the CAP Laboratory Accreditation Program?

Yes

Then your lab must enroll in gene-specific PT such as the MGL2* or MGL5* Survey

Stop

No

Then your lab may enroll in gene-specific PT (such as PGX or CMSP) to satisfy alternative assessment requirements. Otherwise, an alternative assessment scheme, approved by the laboratory director, must be performed

Additional gene-specific PT programs: AAT*, APOE*, BRCA*, CMSP, HGM*, ICSP, IMD*, MGL1-5*, PGX, RETT*, and TPM*

Does your lab do NGS panels, which include reporting/interpreting of specific genes (eg, CFTR associated mutations as part of a large NGS carrier panel)?

Yes

Your lab may enroll in the methods-based NGS-Germline program or NGSE for undiagnosed disorders to satisfy alternative assessment requirements

No

Does your lab do Sanger sequencing on specific genes (eg, cystic fibrosis CFTR screening)?

Yes

If gene-specific PT is not available or not required, your lab may enroll in a methods-based Sanger sequencing (SEC or SEC1) program to satisfy alternative assessment requirements

No

Does your lab do whole genome or exome sequencing via NGS?

Yes

Your lab may enroll in the methods-based NGS-Germline program or NGSE for undiagnosed disorders to satisfy alternative assessment requirements

Stop

No

Your lab may enroll in the methods-based NGS-Germline program or NGSE for undiagnosed disorders to satisfy alternative assessment requirements

A panel is defined as the reporting/interpreting of specific genes on a consistent, ongoing basis, regardless of technical approach (eg, performing exome sequencing on a preselect group of genes would be considered a panel). NOTE: A panel may contain required and non-required analytes; labs must enroll in gene-specific PT for required analytes and may use current PT programs to satisfy alternative assessment requirements for non-required analytes.

*CAP Accreditation Program required program/analyte. Any program without an asterisk (*) reflected in this flow chart is not a required PT program and may be used to fulfill alternative assessment requirements.
Germline Molecular FAQs

Q: My laboratory performs a hearing loss panel by NGS in which we report findings for 100 genes, including GJB2 (Connexin 26). What PT should I enroll in?
A: Your laboratory must enroll in gene-specific PT for Connexin 26 (MGL3* Survey) if it is accredited by the CAP. If there is no gene-specific PT for the remaining genes, your laboratory may enroll in the NGS-Germline program to satisfy alternative assessment requirements. Participation in MGL3* for Connexin 26 (GJB2 gene) will not satisfy alternative assessment requirements for the entire hearing loss panel.

Q: My laboratory tests for rare disorders (eg, Aarskog-Scott syndrome, Von Hippel-Lindau syndrome) by sequencing. What CAP PT is available to satisfy alternative assessment requirements for this assay?
A: Since there is not gene-specific PT available, your laboratory can enroll in the SEC or SEC1 Survey to satisfy alternative assessment requirements for Sanger sequencing and the NGS-Germline Survey to satisfy alternative assessment requirements for NGS. All three Surveys are considered methods-based programs.

Q: My laboratory does exome sequencing on diagnostic odyssey specimens. We report pathogenic/likely pathogenic and variants of uncertain significance that are present in any gene that fits the phenotype. What CAP PT is available to satisfy alternative assessment requirements for this assay?
A: In a case like this, laboratories may enroll in the NGSE Survey to satisfy alternative assessment requirements.

*CAP Accreditation Program required program/analyte.
General Molecular Oncology Flow Chart

Start

Does your lab perform molecular testing on PET?

Yes

Does your lab want challenges in DNA extraction of PET?

Order MHOS

No

Stop

No

Does your lab perform molecular testing on hematologic malignancies?

Yes

Refer to "Hematologic Malignancy Flow Chart"

No

Does your lab perform molecular testing on solid tumors?

Yes

Refer to "Solid Tumor Flow Chart"

No

Does your lab want challenges in tumor cellularity by H&E?

Order NEO

Yes

No

Stop

The testing reflected in this flow chart requires alternative performance assessment. The PT programs listed are not required but may be used to fulfill alternative assessment requirements.
**NOTE:** If there are alternative assays used to confirm NGS findings, these should also have appropriate alternative assessment performed. Alternative assessment options can include participation in CAP PT programs.

*BRAF, KRAS, and EGFR* are CAP Accreditation Program required analytes. Labs must enroll in the MTP Survey or the individual gene Survey programs (EGFR, KRAS and BRAF). Any program without an asterisk (*) reflected in this flow chart is not a required PT program and may be used to fulfill alternative assessment requirements.
Does your lab perform DNA testing for IGH, IGK, TRB, or TRG rearrangements; or IGH/BCL2 or IGH/CCND1 translocations?

- Yes: Order MHO or MHO1*
- No: Does your lab perform qualitative and/or quantitative testing for BCR/ABL1?
  - Yes: p210: order MRD p190: order MRD1
  - No: Does your lab perform quantitative testing for PML/RARA?
    - Yes: Order MRD2
    - No: Does your lab perform qualitative testing for CEBFB/MYH11, RUNX1/RUNX1T1, PML/RARA, BCR/ABL1, CALR, MLL-PTD and other mutations (JAK2 V617F, FLT3 ITD and/or FLT3 TKD, NPM1)?
      - Yes: Order MHO2 or MHO3*
      - No: Order NGSHM

Contains specimens in duplicate.

NOTE: If there are alternative assays used to confirm NGS findings, these should also have appropriate alternative assessment performed. Alternative assessment options can include participation in CAP PT programs.

*The testing reflected in this flow chart requires alternative performance assessment. The PT programs listed are not required but may be used to fulfill alternative assessment requirements.
Molecular Oncology FAQs

Q: My laboratory performs a 50 gene NGS-based assay designed to detect somatic SNVs and small indels observed in solid tumors. What CAP PT is available to satisfy alternative assessment requirements for this assay?
A: BRAF, KRAS, and EGFR are CAP Accreditation Program required analytes and laboratories must enroll in the MTP Survey or the individual gene Survey programs (EGFR, KRAS or BRAF), regardless of the methodology used. The laboratory may enroll in the NGSST Survey to satisfy alternative assessment requirements for the remaining genes in their NGS-based assay.

Q: My laboratory performs a 50 gene NGS-based assay designed to detect somatic SNVs and small indels observed in solid tumors. In addition, we have individual Sanger sequencing-based assays for KRAS, BRAF, and EGFR. Can we use the NGSST Survey to satisfy alternative assessment requirements for all of these analytes/genes?
A: In this case, the laboratory may order the NGSST Survey for their NGS-based solid tumor assay to satisfy alternative assessment requirements and must order either MTP or the individual gene Surveys for KRAS, BRAF, and EGFR.

Q: My laboratory performs a 50 gene NGS-based assay designed to detect somatic SNVs and small indels observed in hematologic malignancies. What CAP PT is available to satisfy alternative assessment requirements for this assay?
A: The laboratory may enroll in the NGSHM Survey. It is not necessary to also enroll in MHO Survey for this assay.
Molecular Oncology FAQs (continued)

Q: My laboratory performs a 50 gene NGS-based assay designed to detect somatic SNVs and small indels observed in hematologic malignancies. In addition, we have individual PCR-based assays for \textit{JAK2}, \textit{FLT3}, and \textit{NPM1}. Can we use the NGSHM Survey to satisfy alternative assessment requirements for all of these analytes/genes?
A: In this case, the laboratory may order the NGSHM Survey for their NGS-based hematologic malignancy assay and MHO2 or MHO3 for the individual PCR-based assays if your laboratory chooses to use CAP PT programs to satisfy alternative assessment requirements.

Q: Our laboratory performs NGS-based testing for the detection of somatic copy number variants and structural variants in solid tumors. What CAP PT is available to satisfy alternative assessment requirements for these assays?
A: Currently, there are no CAP Survey programs for NGS-based detection of copy number variants and structural variants. An alternative assessment scheme, approved by the laboratory director, must be performed (Sample Exchange Registry, etc).

Q: Our laboratory performs NGS-based testing and would like additional bioinformatic challenges in addition to wet-bench challenges. Is there a PT program available for this that may be used to satisfy alternative assessment requirements?
A: Yes, the laboratory may enroll in either NGSB1 if they are using the Illumina TruSeq Amplicon Cancer panel or NGSB2 if they are using the Ion Torrent AmpliSeq Cancer Hotspot v2 panel. At this time, we do not suggest that laboratories performing other panel-based tests use these Surveys. Additionally, there is a somatic validated materials (NGSBV) program available. This in silico program is designed to optimize bioinformatics pipelines, augment validations, and assist with pipeline verification after changes to NGS/bioinformatics processes. This is not traditional PT and no results will be returned to the CAP; information regarding the variants introduced will be sent along with the mutagenized file.
**ISH/FISH Flow Chart**

**Start**

1. **Does your lab test FFPE samples?**
   - Yes
   - **Does your lab test non-FFPE samples (fixed cell suspensions^)?**
     - Yes
     - **HER2 gene Amplification?**
       - Yes
       - Order **ISH***
         - **Kappa/ Lambda, EBV, and/or HPV?**
           - Yes
           - Order **CYJ**
             - **Glioma (1p/19q)?**
               - Yes
               - Order **CYK**
                 - **Solid Tumor?**
                   - Yes
                   - Order **CYL**
                     - **Lymphoma?**
                       - Yes
                       - Order **Cyl**
             - **Glioma (1p/19q)?**
               - Yes
               - Order **CYL**
     - **Brightfield**
       - Yes
       - Order **ISH2***
         - **Kappa/ Lambda, EBV, and/or HPV?**
           - Yes
           - Order **CYH***
             - **Glioma (1p/19q)?**
               - Yes
               - Order **CYJ**
             - **Glioma (1p/19q)?**
               - Yes
               - Order **CYK**
     - **FISH**
       - Yes
       - Order **CYI***
         - **Constitutional/ neoplastic disorders?**
           - Yes
           - Order **CYF**

**Stop**

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*CAP Accreditation Program required program/analyte. Any program without an asterisk (*) reflected in this flow chart is not a required PT program and may be used to fulfill alternative assessment requirements.

**Challenges rotate between sarcomas, neuroblastomas, gastric carcinoma (HER2), and lung (ALK) cancer/carcinoma.**

^ Other preparations (touch preparations, smears, etc.) – alternative assessment required.

ISH/FISH interpretation only – alternative assessment required (split sample exchange, etc.).
Microarray Testing Flow Chart

Start

Does your lab do cytogenomic microarray analysis?

Yes

 Does your lab do expression arrays?

Yes

Constitutional testing?

Yes

Oncology testing?

Yes

Order CYCGH

Order CYCMA

Stop

An alternative assessment scheme, approved by the laboratory director, must be performed

CYCGH PT is not applicable to preimplantation genetic diagnosis (PGD) or exon-level array testing. For PGD, alternative assessment is required. For exon-level arrays, gene-specific duplication/deletion PT may be available (eg, DMD, MECP2) to fulfill alternative assessment requirements or laboratories must identify another form of alternative assessment.

The testing reflected in this flow chart requires alternative performance assessment. The PT programs listed are not required but may be used to fulfill alternative assessment requirements.
Additional Information for Microbiology and Histocompatibility:

Microbiology:
• If performing patient testing on specimens by molecular methods only, labs must meet the regulatory requirements of testing 5 specimens in three mailings for each subspecialty, as appropriate. Subspecialties include Bacteriology, Mycology, Virology, and Parasitology. The Mycobacteriology requirement is 5 specimens tested in each of the two mailings.

• If performing molecular testing on patient specimens, in addition to traditional culture methods, alternative assessment is required. Alternative assessment can be met through enrollment in PT Surveys.

Histocompatibility:
• Regardless of methodology, laboratories should enroll in the appropriate HLA program(s) to meet testing needs.