

should not be more than 10% smaller than the target amplicon and the use of a smaller internal control should be justified.

## BAP.06630 Melting Temperature

Phase I



**For tests that generate a result based on a  $T_m$ , appropriately narrow temperature ranges (+/- 2.5 °C) are defined and recorded each day of use.**

## IN SITU HYBRIDIZATION (ISH)

The use of the term *in situ hybridization (ISH)* in this section applies to all ISH methods, including fluorescence (FISH), chromogenic (CISH), silver (SISH), and brightfield (BRISH) *in situ hybridization*.

Please refer to the Definition of Terms section in the All Common (COM) Checklist for definitions of analytical validation and analytical verification.

### Inspector Instructions:

	<ul style="list-style-type: none"> <li>• Sampling of ISH policies and procedures</li> <li>• Sampling of probe validation/verification records</li> <li>• Sampling of QC records</li> <li>• Sampling of patient test reports</li> </ul>
	<ul style="list-style-type: none"> <li>• How are ISH cut-off values established?</li> <li>• How does your laboratory validate/verify assay performance prior to test implementation?</li> <li>• What is your course of action when a probe does not produce an internal control signal?</li> </ul>

## BAP.06710 ISH Probe Validation/Verification

Phase II

**All in situ hybridization (ISH) probes are validated/verified.**

*NOTE: Additional requirements for test method validation/verification are in the All Common Checklist.*

### Evidence of Compliance:

- ✓ Records of ISH probe validation/verification

### REFERENCES

- 1) American College of Medical Genetics, Standards and Guidelines for Clinical Genetics Laboratories, 2021 edition.
- 2) Clinical and Laboratory Standards Institute (CLSI). *Fluorescence In Situ Hybridization Methods for Clinical Laboratories; Approved Guideline—Second Edition*. CLSI document MM07-A2 (ISBN 1-56238-885-1) Clinical and Laboratory Standards Institute, 940 West Valley Road, Suite 2500, Wayne, Pennsylvania 19087-1898 USA, 2013.
- 3) Lawrence Jennings, Vivian M. Van Deerlin, Margaret L. Gulley (2009) Recommended Principles and Practices for Validating Clinical Molecular Pathology Tests. *Archives of Pathology & Laboratory Medicine*: Vol. 133, No. 5, pp. 743-755
- 4) Wiktor AE, Van Dyke DL, Stupca PJ, et al. Preclinical validation of fluorescence in situ hybridization assays for clinical practice. *Genetics in Medicine* 8:16-23, 2006.
- 5) Weremowicz S, Sandstrom DJ, Morton CC, Miron PM. Validation of DNA probes for preimplantation genetic diagnosis (PGD) by fluorescence in situ hybridization (FISH) R1. *Prenat Diagn.* 2006 Nov;26(11):1042-50.
- 6) Saxe DF, Persons DL, Wolff DJ, Theil, KS; Cytogenetics Resource Committee of the College of American Pathologists. Validation of fluorescence in situ hybridization using an analyte-specific reagent for detection of abnormalities involving the mixed lineage leukemia gene. *Arch Pathol Lab Med.* 2012; 138(1):47-52.

## BAP.06720 Interphase ISH - Cut-off Value

Phase II

**For interphase in situ hybridization (ISH), the biorepository establishes a normal cut-off value for results for each probe used, when applicable.**