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Evolution of Quality Assurance for Clinical Immunohistochemistry in the Era of Precision Medicine: Part 1: Fit-for-Purpose Approach to Classification of Clinical Immunohistochemistry Biomarkers

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From the International Society for Immunohistochemistry and Molecular Morphology (ISIMM) and International Quality Network for Pathology (IQN Path)

Abstract: Technical progress in immunohistochemistry (IHC) as well as the increased utility of IHC for biomarker testing in precision medicine avails us of the opportunity to reassess clinical IHC as a laboratory test and its proper characterization as a special type of immunoassay. IHC, as used in current clinical applications, is a descriptive, qualitative, cell-based, usually nonlinear, in situ protein immunoassay, for which the readout of the results is principally performed by pathologists rather than by the instruments on which the immunoassay is performed. This modus operandi is in contrast to other assays where the instrument also performs the readout of the test result (eg, nephelometry readers, mass spectrometry readers, etc.). The readouts (results) of IHC tests are used either by pathologists for diagnostic purposes or by treating physicians (eg, oncologists) for patient management decisions, the need for further testing, or follow-up. This paper highlights the distinction between the original purpose for which an IHC test is developed and its subsequent clinical uses, as well as the role of pathologists in the analytical and postanalytical phases of IHC testing. This paper is the first of a 4-part series, under the general title of “Evolution of Quality Assurance for Clinical Immunohistochemistry in the Era of Precision Medicine.”

Key Words: biomarkers, quality assurance, quality control, validation, immunohistochemistry

In the era of precision medicine, biomarker testing using immunohistochemistry (IHC) has not only become more precise but also more complex.1–6 Precision medicine requires precision results, which can only come about from precision testing. Because of increasing reliance on
the results of biomarker testing in patient management, a clear understanding by the pathologist of the tests themselves that provide this information is critical to appropriate patient care and thus patient safety.7–9

In this paper, we emphasize the essential role of “purpose” in the IHC assay:
- that “purpose” is the intended use of an IHC test at the time that the test was developed,
- that “purpose” is intrinsic to the identity of any particular IHC test, and
- that classification based on “purpose” of the IHC test is helpful in developing proper quality assurance tools and clarifies the utilities of that test in patient management in the era of precision medicine.10

IHC AS A QUALITATIVE ASSAY FOR THE IN SITU DETECTION OF PROTEIN BIOMARKERS

It is essential to recognize that in its most common application, IHC is a descriptive, threshold-based test. In this context it is generally unknown to what extent the relationship between the amount of target protein and observed intensity of achieved signal is linear.11–14 The intention is to deliver a “positive” or “negative” signal where appropriate and meaningful, rather than to measure the amount of target protein. This mode of application is similar to other testing methods that employ amplification such as polymerase chain reaction (PCR)-based methods, which are also considered as descriptive or “threshold methodologies.” For example, the intention of performing PCR testing is to demonstrate whether DNA or RNA sequences of interest are detected or not, while in IHC the focus is on whether a target protein-based biomarker is detected or not; however, both need to be demonstrated at clinically relevant sensitivity and specificity, which is achieved through validation.15

Today the results of IHC testing are increasingly applied as biomarkers. The term “biomarker” (or biological marker) is broadly defined as any biological or physiological moiety that is used to identify disease, guide targeted therapy or monitor for reoccurrence.16,17 Although the nomenclature is not fully standardized, the National Institutes of Health working group’s definition of a biomarker is more rigid and requires that a biomarker can be objectively measured and evaluated as an indicator of normal biological processes, pathogenic processes, or pharmacologic response to a therapeutic intervention.18 The American Association of Pharmaceutical Scientists and the US Clinical Ligand Society have identified 4 general classes of biomarker assays: (i) definite quantitative assays (eg, mass spectrometry; reference standard is well defined and fully representative of the endogenous biomarker); (ii) relative quantitative assays (eg, ligand-binding assay; reference standards may not be available in a defined form or fully representative of an endogenous biomarker); (iii) quasiquantitative assays (eg, qRT-PCR, which does not employ the use of a calibration standard, but has a continuous response and the result is expressed in terms of a characteristic of the test sample, and (iv) qualitative assays (eg, PCR or IHC; these tests generate categorical data that lack proportionality to the amount of analyte in a sample).8 Qualitative assays such as PCR or IHC typically generate nominal results (eg, presence or absence of target analyte; eg, PCR test positive for Mycobacterium tuberculosis or IHC test positive for S-100 or anaplastic lymphoma kinase (ALK)).8 The results of qualitative assays may also be expressed in ordinal form by use of scoring scales for readout (eg, 0 to 3+ for HER2 IHC in breast cancer). Some IHC tests may have complex readout rules and require either percentage estimate or direct cell counting with predefined cutoff points for what is considered a “positive” or a “negative” result. An example of this approach is the PD-L1 IHC 22C3 PharmDx assay (Dako/Agilent Technologies, Canada) where the cutoff point for a positive test is 50% positive tumor cells.19 Although the percentage of positive cells is reported, no measurement is involved as the overall result is based on identifying the threshold between a positive cell (any membranous staining of tumor cells) versus a negative cell (no staining of tumor cell membrane). Ultimately, therefore, this biomarker is descriptive and qualitative, rather than quantitative in a strict sense.

The future development of IHC and related methods for protein quantification, in the sense of measurement of in situ protein, is beyond the purview this paper and functional capability of IHC as currently practiced. However, the qualitative nature of IHC, as used today in clinical practice, does not make IHC biomarkers inferior to other biomarkers. IHC biomarkers are useful diagnostic, prognostic, and predictive markers when properly validated for specific use.20 Applying identical logic, the PCR test is a valid biomarker for detection of M. tuberculosis, or B or T cell clonality, if the test is properly validated for the specified use.21,22 What ultimately makes a laboratory test “bad” or “good” does not necessarily depend on the level of quantitation in methodology nor on the sophistication of the methodology, but rather, on whether or not the test is properly validated (clinically, diagnostically, and technically) for the specified purpose (“fit-for-purpose”).10

“PURPOSE” IS INTRINSIC TO THE IHC TEST

“Purpose” is intrinsic to the identity of every laboratory test, including IHC tests.8,10,23,24 Because of increasing awareness of the need for quality assurance in every aspect of IHC testing, it is important that the development of each IHC-based assay takes into consideration its intended use at the time of its development. This is particularly relevant to the validation of IHC tests. As we will show in this series, the type of validation depends entirely on the purpose for which a test is developed.10,15,25–27 When the purpose of a test is changed, even if it identifies the same target molecule, it becomes a different test and therefore its validation requirements should be revisited. This can be illustrated by considering the use of antibodies to CD34. Table 1 describes 3 different purposes for performing a CD34 IHC test in clinical practice. As there are at least 3 different purposes for identifying the
CD34 molecule, this in fact represents 3 different CD34 tests, each with its own purpose, test performance characteristics, validation requirements and scoring/readout criteria; this holds true even if for some (different) purposes the CD34 IHC protocol remains the same (Fig. 1).28–31

The question of whether there is such a thing as a “multipurpose” IHC test inevitably arises and it is an important question. The answer is that there can be no such thing as a “multipurpose IHC test” for the reason that purpose is intrinsic to the very definition of any laboratory test including all IHC tests. However, an IHC protocol may certainly be multipurpose, where a single set of protocol conditions may be applied for different IHC tests. For example, it is possible that a single IHC protocol can be used for the purpose of detecting PAX8 expression in tumors of thyroid, kidney, and Müllerian origin. However, it is also possible that an IHC protocol designed to detect PAX8 expression in thyroid tumors, where expression levels are usually high, may fail to detect expression of PAX8 in some other tumors with possibly lower expression levels. Therefore, in the development of this hypothetical PAX8 IHC assay, we would need to consider whether the assay’s analytical sensitivity is fit for each of our various purposes. Although in current clinical practice the majority of IHC protocols are performed under the assumption that they are “multipurpose,” it is unlikely that many such protocols were actually validated for all purposes for which they are being used. Data from proficiency testing (NordiQC, UK NEQAS, etc.) clearly indicates that validation of “multipurpose” IHC protocol is challenging and not uniformly performed by all laboratories. For example, for PAX8, run 42 2014 in the NordiQC program, 25% of the participating laboratories were able to demonstrate PAX8 in the high level expressing sample but produced a false negative result in the sample with low level expression such as clear cell renal cell carcinoma.32

It is of utmost importance that should any new purpose arise for which an existing IHC protocol will be used, it should be carefully re-evaluated and the need for additional validation carefully considered.

**TABLE 1. Different Purposes of the CD34 Immunohistochemistry (IHC) Test**

<table>
<thead>
<tr>
<th>IHC Test</th>
<th>Field</th>
<th>Purpose</th>
<th>Readout</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD34</td>
<td>Dermatopathology and soft tissue pathology</td>
<td>To distinguish solitary fibrous tumor (CD34+) from desmoplastic mesothelioma (CD34-), or To distinguish dermatofibroma (CD34+) from dermatofibrosarcoma protuberans (CD34-)</td>
<td>Positive vs. negative</td>
</tr>
<tr>
<td>CD34</td>
<td>Hematopathology</td>
<td>To identify CD34+ blasts in bone marrow tissue biopsy, to distinguish between acute leukemia (&gt; 20% CD34+ blasts) and myelodysplastic syndrome (MDS) as well as aplastic anemia (no or decreased CD34+ blasts) from hypoplastic MDS (increased CD34+ cells)</td>
<td>Enumeration of CD34+ blasts Counting 500 to 1000 cells Cutoff points at 5% (increased blasts), 10% (accelerated phase of myeloproliferative neoplasms or diagnosis of MDS-II), and &gt; 20% (acute leukemia)</td>
</tr>
<tr>
<td>CD34</td>
<td>Vascular density</td>
<td>To determine degree of vascularization</td>
<td>IA using microvessel density algorithm</td>
</tr>
</tbody>
</table>

IA indicates image analysis.

“**PURPOSE,” “INTENDED USE,” “USE,” AND “FIT-FOR-PURPOSE”**

The “purpose” of an IHC test may be defined as the “intended use at the time the test was developed.” However the “purpose” for which a test was developed and validated by the clinical laboratory may not necessary be the same as the “use” that an ordering physician has in mind (eg, a staff pathologist) after the test has been made available for clinical practice; if such situations arise, then “current use(s)” would no longer align with “intended use at the time the test was developed” (ie, original purpose). For example, the ALK IHC test that was developed and validated for the purpose of detecting anaplastic large cell lymphoma has been “used” to detect ALK expression in non–small cell lung cancer. However, initial attempts resulted in unacceptably low sensitivity because the 2 entities harbor different gene rearrangements and have different protein expression levels; as such, it was discovered that the detection of ALK expression in lung cancer may require an IHC protocol of higher sensitivity than that required for ALK expression in lymphoma.33 In addition, the readout criteria (eg, cutoff values, etc.), which are quite different in lymphomas and non–small cell lung cancer, also must be adapted to reliably predict the clinical relevance.34–36 Therefore, for IHC tests, “purpose” and “intended use at the time the test was developed” reflect test development, which is within the domain of the laboratory. In contrast, “use” in a more general sense relates to clinical practice, which may be different from the original “intended use” and is within the domain of the practicing physician (eg, pathologist and/or treating physician). The concept of “fit-for-purpose” has evolved to address such possible confusion. “Fit-for-purpose” describes an assay that has been successfully validated for the intended use at the time the assay was developed, combining both laboratory and clinical definitions.10,37 It is expected that the new biomarkers will be fit-for-purpose as there are high expectations that the biomarkers will improve diagnosis, define disease subsets that may differ in response, define individual variability in the drug’s molecular target, and provide early clues regarding response to therapy.38,39
From the perspective of the laboratory, purpose is the reason for which the laboratory sets up the test and is what dictates how the laboratory performs the validation of that test. It is therefore important that prior to the development, calibration, and validation of a new IHC assay, there should be coordination between the laboratory director and the end user(s) to ensure that intended use (ie, purpose, which will drive validation requirements) and actual use will be properly aligned. Whether an assay that has been introduced into clinical practice in this manner will ever be “used” for a different purpose than originally intended is unknown to both the laboratory director and to end user(s). However, should such a scenario arise (eg, a different clinical use of an existing assay), a new “fit-for-purpose” validation process would be required.

Lastly, an important point to note regarding “purpose” is that in relation to clinical IHC testing, “purpose” is not simply the in situ detection of a biological gene product in tissue sections. In order to be meaningful, the purpose of any clinical IHC test must always be accompanied by a medical context, because it is in this context that will impact the validation requirements of a particular IHC test.

CLASSIFICATION OF IHC TESTS BASED ON FIT-FOR-PURPOSE PRINCIPLES

A number of classification schemes for diagnostic tests have been introduced in different regions of the world as part of regulatory oversight to ensure appropriate testing for patient care.

The main principle underlying the various classification schemes is based on the degree of risk to patient safety and public health. The degree of oversight for a test should be commensurate with the perceived risk to the patient. The Food and Drug Administration in the United States (FDA) and International Medical Device Regulators Forum have adopted this principle. An approach based on the degree of risk to patient safety translates into the domain of quality assurance, where the test classification dictates differing levels of stringency in relation to requirements for validation, documentation, and reporting. From a quality assurance perspective, approaching the classification of tests in this manner ensures that the “fit-for-purpose” principle is always kept in mind and that the test will be conducted in a manner that considers both the needs of the user and the effect of the test result on the patient.

TYPE 1-IHC AND TYPE 2-IHC TESTS

The Canadian Association of Pathologists (CAP-ACP) has proposed 2 classes of IHC tests based on the end user of the IHC results: class I (pathologist end user) and class II (treating physician end user) IHC tests. The current paper adheres to the same principles but offers an update in terminology in order to highlight and clarify the role of IHC test classification in quality assurance. IHC tests where the test is read by the pathologist, and the results of the readout used by that pathologist for his/her diagnostic practice are termed “pathologist end user IHC tests” (type 1-IHC tests) (Table 2). IHC tests where the pathologist’s readouts are used by a treating physician in order to determine patient management, are “treating physician end user IHC tests” (type 2-IHC tests). The latter include prognostic tests, predictive tests, and screening tests (a special subtype of diagnostic tests) that are relevant for patient management (Table 2).

Figure 1 illustrates similarities and differences between type 1 and type 2 tests.

Although these definitions of type 1-IHC and type 2-IHC tests have broad correlation to various regulatory frameworks, this terminology is introduced to avoid the use of the term “class,” which has been defined on a different basis. The goal is to avoid potential confusion with FDA classification of IHC devices and other test classification schemes in various countries that address risk and refer to industry and devices rather than IHC tests and how these are actually used by pathologists and/or treating physicians. The comparison with classification schemes used by various regulatory agencies is shown in Table 3.

THE IHC PROTOCOL, THE “READOUT” AND THE "INTERPRETATION"

The IHC test encompasses preanalytic, analytic, and postanalytic phases. Figure 1 illustrates these different components of the IHC test. The analytical phase of the IHC test is complex and it consists of various components of the IHC protocol as well as the “readout” of the generated IHC slide. There is an essential difference between IHC testing and other biomarker testing methods. As noted above, IHC tests are for the most part descriptive, or at best quasiquantitative, and the readout is the culmination of a morphologic analysis of an IHC slide by a pathologist, who assesses the presence and localization of signals, evaluates cutoff points between what
constitutes a positive and negative signal, and counts positive cells, or uses other methods of scoring as appropriate. In essence, the IHC test is always descriptive, even when it is possible to provide information in a quantitative format.

It takes years of training and experience as well as focused continuing medical education to become proficient at reading IHC slides. For certain biomarkers, the readout criteria to determine whether a test is “positive” or “negative” may be seemingly straightforward (e.g., reading thyroid transcription factor-1 IHC slides in lung cancer); however, it may also be very complex such as when multiple criteria are incorporated in the readout (e.g., reading HER2 IHC slides) or when evaluating for vascular invasion, microinvasion, micrometastasis, where higher levels of experience are required for proper readout. This practice is clearly in contrast to other biomarkers for which readouts are performed by automated instruments of different types (nephelometry, colorimetric methods, mass spectrometry, flow cytometry, etc.). Computer-aided digital image analysis (IA) has been introduced for readout of IHC slides, with the intent of improving reproducibility of results. IA currently still requires time from a pathologist to determine what part of the tissue and what cells will be scored as well as which algorithms are applicable for a given readout; a number of these algorithms have been approved by the FDA and other organizations for clinical use. In addition, the approval of whole slide imaging for primary diagnosis, already effected in Europe and Canada, and under active trial in the United States, surely will impact the growth of computer-assisted morphometry and analysis of IHC in the near future. Nevertheless, at present, IA should be viewed as assistive technology that helps, but currently cannot replace, pathologists in generating the readout from an IHC slide. The analytical phase of IHC testing is completed by the readout of the IHC slide.

Interpretation of the readout occurs in the postanalytical phase of IHC testing. For type 1-IHC tests, it is the pathologist end user who interprets the readout results. An example of this would be a clinical case of “tumor of unknown origin” where IHC testing was ordered to help identify the potential primary site of a metastatic tumor deposit. A typical readout result might be that cytokeratin 20 and CDX2 are positive and cytokeratin 7 and thyroid transcription factor-1 are negative;

### TABLE 3. Comparison of Immunohistochemistry (IHC) Classification Schemes of IHC Tests by Regulatory Agency and Classification Based on Clinical Practice

<table>
<thead>
<tr>
<th>Regulation of Test Manufacturers</th>
<th>Intention</th>
<th>Class</th>
<th>Clinical Practice (Nearest Correlate)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA</td>
<td>For regulating manufacturers of tests</td>
<td>Class 1</td>
<td>Type 1-IHC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class 2</td>
<td>Type 2-IHC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class 3</td>
<td>Type 2-IHC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class 1</td>
<td>Type 1-IHC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class 2</td>
<td>Type 2-IHC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class 3</td>
<td>Type 2-IHC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class 4</td>
<td>Type 2-IHC</td>
</tr>
<tr>
<td>Health Canada</td>
<td>For regulating manufacturers of tests</td>
<td>Class A</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class B</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class C</td>
<td>Type 1-IHC, type 2-IHC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class D</td>
<td>Type 2-IHC</td>
</tr>
<tr>
<td>EU In-vitro Diagnostic Regulation</td>
<td>For regulating manufacturers of tests</td>
<td>Class 1</td>
<td>Type 1-IHC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Class 2</td>
<td>Type 2-IHC</td>
</tr>
<tr>
<td>Canadian Association of Pathologists</td>
<td>Guidance for clinical practice</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

FDA indicates Food and Drug Administration in the United States; NA, not available.
the subsequent interpretation by the pathologist end user (ie, the sign-out pathologist) may then reasonably be “consistent with adenocarcinoma with intestinal differentiation.”\textsuperscript{62} The sign-out pathologist in this example may or may not choose to report the detailed readout of the IHC results, but will certainly report his/her interpretation of the readout results. It is also worth noting that for type 1-IHC tests, which by current convention are both “read” and “interpreted” by the same pathologist (ie, the sign-out pathologist), it may appear as though readout and interpretation are concurrent; in practice they may occur very close together, but the interpretation is necessarily subsequent to the readout.

For type 2-IHC tests, it is the treating physician end user who interprets the readout results in the context of clinical actions. An example would be a treating physician who receives IHC results for a patient with carcinoma of the breast stating that the sampled tumor cells are “strongly positive for estrogen receptor and progesterone receptor and negative for HER2”\textsuperscript{2}; the treating physician may then interpret the readout results as indicating that “the patient is a good candidate for hormonal therapy.”\textsuperscript{63}

When a pathologist reports a test result as “positive” versus “negative,” the presumption is that the pathologist is following predetermined scoring criteria using predetermined scoring or cut off points, extending beyond the usual meaning of “interpretation.” As the term “interpretation” is inherently associated with postanalytical phase, it is important that any readout as a part of analytical phase, irrespective of its complexity, is not called “interpretation.” Moreover, in some type 2 tests, pathologists may also contribute more directly to the clinical interpretation that will eventually help the treating physician in patient management. For example in IHC testing for DNA mismatch repair deficiency, pathologists may provide additional interpretation of unusual patterns (eg, when reporting discordant subclonal patterns) that directly impact upon how the treating physician interprets and applies the results in treatment.\textsuperscript{64}

This paper seeks to apply the standardized terminology for laboratory testing in the context of IHC; thus “readout” is considered to be part of the analytic phase, whereas “interpretation” is linked to the postanalytical phase.\textsuperscript{65} For clarity, the term “readout” is applied to the process whereby a pathologist using high-level skills “reads” the IHC slide (ie, produces a result), before a closely related but subsequent process in which the readout (or IHC result) is interpreted in the context of pathology diagnosis or clinical scenario. Clearly, pathologists are “interpreting” (in the more general use of the word) all visual cues present in the IHC slides in arriving at the most accurate readout (result) (eg, assessing signal intensity, localization, “texture,” and other signal parameters in conjunction with controls, excluding various type of artifacts, etc.). However, all subsequent mental processing of the information incorporated into the “readout” is, by analogy with other forms of laboratory testing, properly considered to be “interpretation” by the pathologist (for type 1 IHC results) for diagnostic purposes or by the treating physician (for type 2 IHC results) for decisions directly relevant to patient treatment. With this approach, the readout is properly identified as a part of the analytical phase, just like it is in any other laboratory test (eg, nephelometry, mass spectrometry, etc.).\textsuperscript{66,67} Furthermore, this approach also applies to IA; IA is intuitively accepted as “readout,” not as “interpretation” because it is performed by machines executing automated algorithms rather than by a living person.

CONCLUSIONS

Biomarker testing by IHC is critical to patient care in the era of precision medicine. The IHC test is purpose-driven and purpose-defined. In today’s world, an IHC test is only meaningful when it is linked with a specific purpose for which it has been developed. Part 1 of this 4-part series highlights the following:

1. The purpose of an IHC test is intrinsic to the test itself; if it is discovered that the current use of the test is different from its original purpose, the new purpose requires consideration of whether or not a new fit-for-purpose development and validation process is needed.

2. The result of an IHC test is not the IHC slide itself with the signal pattern that emerges on the tissue after development of the reaction product; rather, it is the readout of that signal pattern. This readout is performed by a pathologist, sometimes with the assistance of computer aided IA. Readout of IHC slides does require intellectual activity, observation and judgment, but this process should be considered as separate from interpretation of laboratory results in a strict sense, which is about what the result “means.”

3. The readout (test result) is a part of the analytical phase of IHC testing and is distinct from interpretation of the meaning, significance and utility of the test result. Generally, interpretation is a postanalytic test element performed by a pathologist (for diagnosis) or a treating physician (± pathologist) (for treatment decisions).

4. Using a fit-for-purpose approach, IHC tests can be classified as “type 1-IHC” tests or “type 2-IHC” tests based on the end-user.

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