ALSO IN THIS ISSUE:

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Clinical and Translational Pathology Innovations
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Feature Story
Quantification of drugs/metabolites in urine by liquid chromatography-tandem mass spectrometry for pain management services

Molecular Diagnostics in BCR-ABL1 Negative Myeloproliferative Neoplasms | p 6
STAT6 Rabbit Monoclonal Antibody for the Diagnosis of Solitary Fibrous Tumor | p 10
 Alumni Connect | p 15
News | p 16
Quantification of drugs/metabolites in urine by liquid chromatography-tandem mass spectrometry for pain management services

By Dustin Bunch, Katherine Lembright and Sihe Wang, PhD*

Pain management drugs are among the most prescribed medications in the United States and are often abused.1, 2 It is estimated that between 9 to 41% of patients receiving opioids for chronic pain abuse the prescription.3, 4 Chronic pain patients also are estimated to use illicit drugs at a rate of 14 to 34%.2 A serious public health problem is caused by the diversion of prescribed pain management drugs.6

Monitoring the drugs and metabolites of the prescribed medication and potentially abused drugs plays an important role in ensuring patient compliance. Urine is the preferred specimen due to the ease of sample collection and the acquired background knowledge for this sample type. Urine drug testing has often been used to verify adherence to prescribed pain management regimens and to detect illicit drug use.7 Both routine and random drug testing can significantly improve patient compliance.8-10 The test results have important implications regarding the patient’s eligibility for additional pain management medications. Therefore, accuracy in determining both negative and positive results is extremely important.

Though immunoassays are easy to run with fast turnaround time, they may lack the needed sensitivity and/or specificity.11, 12 While gas chromatography-mass spectrometry (GC-MS) methods are considered the gold standard, sample preparation is labor intensive. There is increasing use of liquid chromatography-tandem mass spectrometry (LC-MS/MS), which generally requires much simpler sample preparation with high specificity and sensitivity.13

Some drugs/metabolites, such as morphine and codeine, are present in urine as both free and glucuronide conjugated forms. Percentage of the conjugates may vary significantly depending on both sample collection time and individual metabolism rates.14 Conjugated analytes may be hydrolyzed prior to LC-MS/MS analysis in order to improve sensitivity and consistency. While chemical hydrolysis is fast and efficient, it may result in undesired reactions and loss of important information. For example, acid hydrolysis converts heroin and its metabolite, 6-acetylmorphine, to morphine.15 In contrast, enzymatic hydrolysis is more specific, but requires longer incubation time to achieve high efficiency (> 90%).16

While direct injection of diluted urine samples for LC-MS/MS analysis has been reported, the matrix effect may compromise sensitivity and accuracy.17 Both off-line and online solid phase extraction is widely used for sample purification prior to LC-MS/MS analysis18-21 to reduce the matrix effects. Mueller et al. reported an online turbulent flow extraction method using two TurboFlow columns with different stationary phases to extract hundreds of drugs/metabolites.22 Most of the analytes important for pain management are measured in the positive electrospray ionization (ESI) mode,18-20, 23 while tetrahydrocannabinol carboxylic acid (THCA), the major metabolite of marijuana, is known to ionize more efficiently in negative ESI mode.24

At Cleveland Clinic, we have developed and validated a novel LC-MS/MS method for simultaneously measuring 20 drugs and metabolites in urine to monitor the use of 16 prescription or illicit drugs (morphine, codeine,
dihydrocodeine, oxycodone, oxymorphone, hydrocodone, hydromorphone, methadone, fentanyl, tramadol, buprenorphine, amphetamine, methamphetamine, cocaine, heroin and marijuana. The chromatogram is shown in Figure 1. Sample preparation includes enzymatic hydrolysis followed by online turbulent flow extraction. Recovery for all analytes ranges between 93.2% and 111.4%. The lower limit of quantification (LLOQ) is between 5 and 25 ng/mL for all analytes. Accuracy varies between 85.8% and 119.4%. Intra-assay and total CVs at three different levels are 0.2% – 12.7%.

Significant inter-laboratory variation has been reported when measuring conjugated urine drugs, especially for codeine. The high variation was likely due to different hydrolysis methods used in each laboratory. Enzymatic hydrolysis efficiency is greatly affected by the type of glucuronidase and the reaction condition used. In addition, hydrolysis rates of different drugs vary significantly under the same reaction condition. Therefore, a robust and reproducible hydrolysis method is important for consistent measurement. Compared to other glucuronidases, the glucuronidase from Patella vulgata showed a superior efficiency to liberate morphine from its glucuronide conjugates. We found that reproducible and near complete (> 94%) hydrolysis for all analytes can be achieved using this enzyme with an overnight incubation (≥ 16 h) at 60°C.

Simultaneously measuring a large panel of pain management drugs in one LC-MS/MS run has obvious benefits. However, significant variances in their chemical properties can make concurrent measurement technically challenging. As a result, separate sample preparation and separate MS methods were required to quantify different classes of drugs in the past. We use two TurboFlow columns with compensatory stationary phases to simultaneously extract 20 analytes. These two TurboFlow columns in tandem allow a clinically meaningful LLOQ to be achieved for all analytes. Another advantage of online extraction is reduced labor while achieving a high reproducibility.

In summary, we have established a novel quantitative urine drug analysis to simultaneously monitor the use of 16 drugs. This method has been successfully used to support pain management clinics in the last six years with ~85,000 samples analyzed. This method also has successfully passed multiple CAP proficiency testing challenges.

References:


About the Authors

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Dustin R. Bunch received his bachelor’s degree in biochemistry from Case Western Reserve University and is currently a Senior Research Technologist in the Department of Clinical Pathology and a graduate student in the Clinical and Bioanalytical Chemistry program at Cleveland State University. His major duties involve managing the research team and performing development and validation of liquid chromatography tandem mass spectrometry (LC-MS/MS) methods. He has authored or co-authored more than 50 peer-reviewed abstracts and publications and has been a member of the American Association for Clinical Chemistry (AACC) since 2008. Upon completion of his PhD, Dustin plans to join a Commission on Accreditation in Clinical Chemistry (ComACC) approved post-doctoral fellowship program in clinical chemistry. He currently serves as the Internet Coordinator for the Northeast Ohio Section of AACC and the Ohio Collaborative Laboratory Conference. Mr. Bunch can be reached at bunchd@ccf.org or 216.444.7003.

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Molecular Diagnostics in BCR-ABL1 Negative Myeloproliferative Neoplasms

By David Bosler, MD

Advancements in the understanding of disease and the molecular diagnostics of myeloproliferative neoplasms (MPNs) have significantly changed their diagnosis. Although not as dramatic in terms of effect on disease management as those seen in BCR-ABL1 positive chronic myelogenous leukemia (CML), multiple recent advancements in the diagnosis of BCR-ABL1 negative MPNs make diagnosis of these entities today both simpler and more precise than just ten years ago. This article highlights recent developments in the diagnosis of BCR-ABL1 negative MPNs and their practical impact on the diagnostic algorithm.

A key decision point encountered in the diagnosis of MPNs is the distinction between BCR-ABL1 positive CML and BCR-ABL1 negative MPNs. The testing for diagnosis and management of CML is largely unchanged since the review in the Fall 2012 issue of Pathology Innovations and is not discussed in any detail here. However, it is important to remember that, although BCR-ABL1 positive CML most often has a classic presentation that is recognizable by peripheral blood and marrow cell counts and morphology, CML has a spectrum of morphologic presentations that can overlap with BCR-ABL1 negative MPNs as well as other entities (including PDGFRA, PDGFRB and FGFR1 associated neoplasms and some myelodysplastic/myeloproliferative neoplasms). Exclusion of BCR-ABL1 fusion is therefore an important component of the work-up for these neoplasms. Even though misdiagnosis of this type is a relatively uncommon event, it is important not to miss BCR-ABL1 positive CML given its dramatic response to tyrosine kinase inhibitor therapy.

One of the most important tasks when evaluating a patient for MPNs is the distinction between benign and malignant. The overlap of clinical and morphologic features between the BCR-ABL1 negative MPNs and benign, reactive processes has historically meant that definitive diagnosis relied on a combination of sometimes laborious testing (such as the red blood cell mass) and

<table>
<thead>
<tr>
<th>Test Name</th>
<th>Test Code</th>
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<tbody>
<tr>
<td>JAK2 V617F Mutation Detection</td>
<td>JAK2</td>
</tr>
<tr>
<td>JAK2 Exon 12 - 15 Sequencing</td>
<td>JAKNON</td>
</tr>
<tr>
<td>MPL Mutation Analysis</td>
<td>MPL</td>
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<tr>
<td>CALR (Calreticulin) Exon 9 Mutation Analysis</td>
<td>CALR</td>
</tr>
<tr>
<td>Myeloid Malignancies Mutation Panel by Next Generation Sequencing</td>
<td>MYENGS</td>
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exhaustive exclusion of potential non-neoplastic etiologies. Recently developed molecular diagnostics tools that aid in definitive diagnosis of BCR-ABL1 negative MPNs have greatly simplified the diagnostic process. Until about ten years ago, molecular diagnostics was only of benefit in diagnosing BCR-ABL1 negative MPNs in those few cases (less than 10%) where cytogenetic karyotyping detected an abnormal clone. The discovery of JAK2 mutations in MPNs in 2005 began to change that.

The most prevalent JAK2 mutation in MPNs is V617F, a single nucleotide substitution on chromosome 9p, within an auto-inhibitory domain of JAK2. The mutation results in constitutive activation of the JAK-STAT pathway, driving proliferation and survival even in the absence of growth factor stimulation. Originally described in polycythemia vera (PV), the JAK2 V617F mutation is present in 97% of PV and also in at least half of the two other BCR-ABL1 negative MPNs, essential thrombocythemia (ET) and primary myelofibrosis (PMF). Although the JAK2 V617F mutation can be present in extremely low levels in some individuals without myeloid neoplasia, when positive at the level detected by most clinical assays in the context of the appropriate clinical and pathologic context, the mutation is a strong indicator of myeloid neoplasm.
Since the discovery of JAK2 V617F, MPN clinical diagnostics development has focused on what to do about the cases that are negative for V617F—how can they be diagnosed with the same ease as V617F positive cases? These gaps have been closed over time in piecemeal fashion. Essentially all of the PV cases that are negative for V617F have one of a variety of less common mutations that can be detected by sequencing of JAK2 exons 12-15. Non-V617F mutations are found in PV, but not ET or PMF.

MPL gene mutations are present in small percentages of ET (1-4%) and PMF (5-11%), but not PV. Functionally, these mutations act similarly to the JAK2 mutation, except that the mutation is directly in the thrombopoietin receptor, constitutively activating it rather than a downstream tyrosine kinase as in the JAK2 mutation. The relevant MPL mutations are in exon 10 and the vast majority are single nucleotide substitutions at position 515, making a variety of methodologies viable for detection. Other single nucleotide substitutions and deletion/insertion mutations near position 515 have also been described. MPL mutations add value to the diagnosis of MPNs since they are mutually exclusive with JAK2 mutations and are detected only in myeloid neoplasms (mainly MPNs).

Recently, sequencing studies on JAK2-negative, MPL-negative MPNs led to the discovery of mutations in the calreticulin (CALR) gene in a significant percentage of these cases. The CALR gene is on chromosome 19p13.2 and contains nine exons. CALR mutations are present in up to 90% of JAK2-, MPL- ET and PMF cases. Reported mutations are frame shift mutations within exon 9, with two variants (one 52 bp deletion and one 7 bp insertion) representing more than 80% of mutated cases. Like MPL mutations, CALR mutations are found in ET and PML but not in PV. CALR, JAK2 and MPL mutations are mutually exclusive. CALR mutations are also variably reported to be present in low percentages of myelodysplastic syndromes (MDS) and overlap MDS/MPNs. Although the mechanism of the mutation in pathogenesis is unknown, over-expression of mutated CALR in in vitro models showed cytokine independent growth through STAT5 activation. MPN cases with mutated CALR appear to have comparatively favorable prognosis, although there is currently insufficient data to justify any differences in therapeutic approach.

Combined, the JAK2, MPL and CALR mutations now account for more than 90% of BCR-ABL1 negative MPNs. Access to assays that detect these mutations will result in earlier and more efficient diagnosis or exclusion of disease in suspected cases. Work-up of suspected BCR-ABL1 negative MPN should therefore routinely include these tests. Although the actual testing algorithm used may depend on the availability of specific tests, samples and nucleic acid extraction protocols, a recommended approach is shown in Figure 1. After BCR-ABL1 is excluded, JAK2 V617F has the highest diagnostic yield and should generally be tested first. Although disease burdens vary, the allele-specific PCR method used in JAK2 V617F has an analytic sensitivity of 1%, which is sufficient for detection of JAK2 positive MPNs. If V617F is negative, the next step depends on which disease is suspected and the level of clinical suspicion. If PV is highly suspected despite negative JAK2 V617F (eg. chronic polycythemia and low erythropoietin level), JAK2 sequencing of exons 12-15 should be next. While a positive sequencing result is diagnostic and most positive cases will be detected, a negative result does not completely exclude the possibility of very low levels of PV since Sanger sequencing requires at least 20% mutated allele burden to detect a mutation. This test should not be used for routine screening of erythrocytosis, but should instead be reserved for those cases where clinical suspicion is high and persists even after demonstrated absence of JAK2 V617F.

If the clinical and morphologic evaluation instead suggests ET or PMF, CALR and MPL testing is recommended for
JAK2 V617F negative cases. The most cost-effective way to perform this testing is to evaluate for JAK2, CALR and MPL in sequential order. If testing is positive at any step, no further testing is necessary and a diagnosis can be made based on the marrow morphologic findings. At Cleveland Clinic, CALR mutation testing is performed using PCR with fragment length analysis, which has an analytical sensitivity of approximately 5%. MPL mutation testing is performed by Sanger sequencing of exon 10. This method provides comprehensive detection of the MPL mutations described as well as potential mutations within the region of interest on exon 10. As a Sanger sequencing method, the MPL sequencing assay is subject to the same limitations in detecting very low disease burdens as described for the JAK2 sequencing assay above.

Although each of these tests has some diagnostic value as described above, they have relatively limited use in classifying BCR-ABL1 negative MPNs into one of the specific 2008 WHO diagnoses (PV, ET, PMF). Accurate diagnosis and sub-classification is nonetheless clinically relevant due to prognostic differences as well as some variation in approaches to treatment. Molecular findings can provide some clues such as JAK2 non-V617F indicating PV, and CALR or MPL indicating non-PV, but the specific classification usually relies on clinical and morphologic evaluation.

Cytogenetic karyotyping should also be performed on all bone marrows performed for diagnosis of MPNs. Karyotyping provides a baseline karyotype used to track clonal evolution over time and also provides prognostically relevant information. For the very rare cases where MPN is highly suspected based on the clinical and/or morphologic findings and all testing is negative, a multi-gene sequencing panel containing genes relevant to myeloid neoplasia, such as ASXL1, TET2, DNMT3A, SF3B1, may provide additional information helpful in yielding a definitive diagnosis. Such testing is not required for diagnosis according to 2008 WHO criteria, provided that diagnostic morphologic features are present and non-neoplastic etiologies have been excluded.

Another development in the diagnosis of MPNs is the recent description of mutations in the colony stimulating factor receptor 3 gene (CSF3R) in chronic neutrophilic leukemia (CNL). CNL is a rare disease, representing much less than 1% of MPNs. It can present a diagnostic challenge however, since the morphologic findings are essentially indistinguishable from those of reactive neutrophilia (which is vastly more common) and also due to overlap with the morphologic spectrum of BCR-ABL1 positive CML. Like other MPNs, diagnosis of CNL has historically relied heavily on exhaustive exclusion of other potential etiologies. Until CSF3R mutations were first described in CNL by Maxson et al. in 2013, CNL had lacked a distinctive molecular marker, with almost 90% of cases having a normal cytogenetic karyotype. Located on chromosome 1p34.3, CSF3R encodes for the granulocyte stimulating factor transmembrane receptor, which plays a role in granulocyte growth and differentiation. Mutations in CSF3R have also been described in some congenital neutropenias, and in hereditary neutrophilia. The most common CSF3R mutation in CNL is T618I, which is present in up to 83% of studied cases and appears to show specificity for CNL. CSF3R mutation testing will significantly change the diagnostic algorithm for CNL, and it has been proposed that CSF3R be incorporated as a major diagnostic criterion for the next edition of the WHO classification. Given that reactive neutrophilia is so common and CNL so rare, CSF3R testing should not be used to routinely screen neutrophilic patients. Testing should be limited to cases with high clinical suspicion (chronic, persistent neutrophilia with no evident underlying etiology) and where definitive diagnosis would impact clinical management.

In summary, the advances in molecular diagnostics in the past ten years have transformed the way myeloproliferative neoplasms are diagnosed. Used wisely in conjunction with clinical presentation, select laboratory results and marrow morphology findings, JAK2 V617F, JAK2 sequencing, CALR and MPL assays can be valuable tools in diagnosis of PV, ET and PMF. Although CSF3R is also now a potential tool in diagnosis of CNL, to avoid over-utilization it should only be used when clinical suspicion is very high since CNL is so rare.

References:


About the Author

David Bosler, MD

David Bosler, MD, is Head of Cleveland Clinic Laboratories, Staff in Laboratory Medicine at Cleveland Clinic and Assistant Professor at the Cleveland Clinic Lerner College of Medicine. He is an AP/CP certified pathologist with subspecialty certifications in Hematopathology and Molecular Genetic Pathology. His clinical practice includes bone marrow biopsy interpretation and development and interpretation of molecular hematology tests. He has authored or co-authored numerous peer-reviewed journal articles and book chapters. Dr. Bosler completed medical school at University of Cincinnati, residency at William Beaumont Hospital, and fellowships at Mayo Clinic. He has also previously served as Medical Director for Point of Care Testing and Chair of the Point of Care Compliance Council at Cleveland Clinic. Dr. Bosler can be reached at boslerd@ccf.org or 216.636.9615.
The STAT6 rabbit monoclonal antibody is an extremely sensitive and specific marker for the diagnosis of solitary fibrous tumor, a fibroblastic tumor of intermediate biological potential that can arise in a variety of anatomical sites, including pleura, meninges, extrapleural soft tissue and viscera.\(^1\)

Solitary fibrous tumor is classically characterized by CD34-positive spindle cells in a collagenous background with an elaborate vasculature that includes staghorn blood vessels. However, the morphological spectrum of solitary fibrous tumor is broad and encompasses cellular variants (formerly known as hemangiopericytoma), fat-forming variants (also known as lipomatous hemangiopericytoma), giant cell angiofibroma and malignant solitary fibrous tumor.\(^2-10\) This morphological spectrum raises a broad differential diagnosis. Many entities that are included in the differential diagnosis show striking morphological resemblance and share CD34 expression. Approximately 5–10% of solitary fibrous tumors are negative for CD34 and, in this setting, the diagnosis can be especially challenging.\(^1,2,11,12\)

Most solitary fibrous tumors are clinically indolent; however, approximately 10% behave more aggressively with local recurrences and metastases. The mainstay of therapy is wide en bloc resection with indefinite follow up.\(^1\)

Recent studies using next generation sequencing techniques demonstrated the presence of recurrent fusions between NAB2 and STAT6 on chromosome 12q13 in the majority of solitary fibrous tumors,\(^13-15\) with a \(\text{NAB2-STAT6}\) fusion transcript detected in 55–100% of tumors, regardless of tumor morphology (benign, lipomatous, malignant) or anatomical site.\(^13-15\) Schweizer et al. recently investigated immunohistochemistry for STAT6 as a surrogate marker of the \(\text{NAB2-STAT6}\) fusion in meningeal hemangiopericytoma/solitary fibrous tumors.\(^16\) The \(\text{NAB2-STAT6}\) fusion leads to EGR1 activation and transcriptional deregulation of EGR1-dependent target genes and is a driving event in initiation of solid fibrous tumors.

### Table 1. STAT6 expression in solitary fibrous tumor by histological subtype

<table>
<thead>
<tr>
<th>Histological subtype</th>
<th>STAT6 nuclear staining, positive cases/total cases (%)</th>
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<tbody>
<tr>
<td>Fibrous</td>
<td>23/23 (100%)</td>
</tr>
<tr>
<td>Cellular (hemangiopericytoma)</td>
<td>18/18 (100%)</td>
</tr>
<tr>
<td>Giant cell angiofibroma</td>
<td>2/2 (100%)</td>
</tr>
<tr>
<td>Malignant/dedifferentiated</td>
<td>11/11 (100%)</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>54/54 (100%)</strong></td>
</tr>
</tbody>
</table>

### Table 2. STAT6 expression in histological mimics of solitary fibrous tumor

<table>
<thead>
<tr>
<th>Soft tissue tumor</th>
<th>STAT6 nuclear staining, positive cases/total cases (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spindle Cell Lipoma</td>
<td>cell lipoma 0/10 (0%)</td>
</tr>
<tr>
<td>Mammary-type myofibroblastoma</td>
<td>0/10 (0%)</td>
</tr>
<tr>
<td>Cellular angiofibroma</td>
<td>0/9 (0%)</td>
</tr>
<tr>
<td>Benign fibrous histiocytoma</td>
<td>0/13 (0%)</td>
</tr>
<tr>
<td>Dermatofibrosarcoma protruberans</td>
<td>0/9 (0%)</td>
</tr>
<tr>
<td>Desmoid-type fibromatosis</td>
<td>0/8 (0%)</td>
</tr>
<tr>
<td>Low grade fibromyxoid sarcoma</td>
<td>0/7 (0%)</td>
</tr>
<tr>
<td>Schwannoma</td>
<td>0/8 (0%)</td>
</tr>
<tr>
<td>Malignant peripheral nerve sheath tumor</td>
<td>0/7 (0%)</td>
</tr>
<tr>
<td>Monophasic synovial sarcoma</td>
<td>0/11 (0%)</td>
</tr>
<tr>
<td>Mesenchymal chondrosarcoma</td>
<td>0/7 (0%)</td>
</tr>
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STAT proteins are transcription factors that are normally located in the cytoplasm in latent form and migrate to the nucleus after cytokine exposure and subsequent phosphorylation. In cells harboring a NAB2-STAT6 fusion, STAT6 is similarly relocated to the nucleus.

Nuclear expression of STAT6 was detected by immunohistochemistry in 35 of 37 meningeal hemangiopericytoma and 25 of 25 meningeal solitary fibrous tumors, supporting the idea that meningeal hemangiopericytoma and meningeal solitary fibrous tumor are part of the same histological spectrum of solitary fibrous tumor and not distinct entities. Nuclear expression of STAT6 also discriminated solitary fibrous tumor from its various morphological mimics in the meninges, including meningioma, glioblastoma, gliosarcoma, hemangioblastoma, schwannoma and hemangioma.

Our study examined the sensitivity and specificity of STAT6 immunohistochemistry in reliably distinguishing solitary fibrous tumors from other soft tissue neoplasms that are in the differential diagnosis. We used a STAT6 rabbit monoclonal antibody (1:100), which had not previously been reported by others, on formalin-fixed, paraffin-embedded whole sections and tissue microarray slides. STAT6 expression was evaluated in 54 solitary fibrous tumors from 26 males and 28 females and 99 soft tissue tumors in the histological differential diagnosis. Only nuclear staining of STAT6 was considered positive. Distribution of staining was scored as: 0 (no staining), 1+ (1-25%), 2+ (26-50%), 3+ (>50%). Intensity was scored as weak, moderate or strong. Nuclear STAT6 staining was present in all SFT cases tested (54/54, sensitivity 100%), regardless of histology, anatomical site or CD34 status. The majority of cases showed 3+ and strong staining. All tested cases of cellular angiofibroma (0/9), myofibrosarcoma (0/13), dermofibrosarcoma protuberans (0/9), low-grade fibromyxoid sarcoma (0/7), schwannoma (0/8), desmoid-type fibromatosis (0/8), monophasic synovial sarcoma (0/11), malignant peripheral nerve sheath tumour (0/7), and mesenchymal chondrosarcoma (0/7) were negative for STAT6 (specificity 100%).

The study further supports the utility of STAT6 immunohistochemistry as an adjunct in the diagnosis of solitary fibrous tumor.

The study demonstrated that the STAT6 rabbit monoclonal antibody is an extremely sensitive and specific marker for the diagnosis of solitary fibrous tumor, with 100% sensitivity and 100% specificity (no dedifferentiated liposarcomas were tested in this study). Positive nuclear staining for STAT6 was observed in all 54 solitary fibrous tumors, regardless of histology, anatomical site or CD34 status. The results are in concordance with previously published studies of STAT6, but differ from those studies in the antibody that was used. During the period of our validation study, Doyle et al. reported 98% sensitivity in a similar study of 59 solitary fibrous tumors versus other spindle cell neoplasms using a STAT6 rabbit polyclonal antibody.

In our experience, the STAT6 rabbit monoclonal antibody showed clean nuclear staining with little to no background, and we observed no cytoplasmic staining; consequently, all of our negative cases were easily interpretable and we did not encounter any “false positive” cases.

In practice, the diagnosis of solitary fibrous tumor is usually straightforward; however, occasionally it can present as a diagnostic challenge, for example, in the setting of CD34 negative solitary fibrous tumors and malignant/ dedifferentiated solitary fibrous tumors. Much emphasis has traditionally rested on the expression of CD34 in the diagnosis of solitary fibrous tumor, which supports the diagnosis, but is also shared by several other soft tissue tumors that show morphological similarity, including spindle cell lipoma, mammary-type myofibroblastoma, cellular angiofibroma and dermofibrosarcoma protuberans. The absence of CD34 expression also does not exclude the diagnosis of solitary fibrous tumor, since a small subset of solitary fibrous tumors is CD34 negative. Other markers such as BCL2 and CD99 are variably used to support the diagnosis of solitary fibrous tumor but are less sensitive than CD34 and equally non-specific. In diagnostically challenging cases, particularly on core needle biopsies, immunohistochemistry for STAT6 can be a valuable diagnostic adjunct.
Ongoing advances in our understanding of the biology of solitary fibrous tumor lends itself to further work on pathway-specific targeted therapies for inoperable or malignant disease.

References:


About the Author

**Brian B. Rubin, MD, PhD**

Brian B. Rubin, MD, PhD, is Professor of Pathology and Vice-Chair of Research at the Robert J. Tomsich Pathology & Laboratory Medicine Institute. He is a world-renowned molecular pathologist and researcher with a successful track record and a focused interest in the diagnosis and treatment of sarcomas. Dr. Rubin is author of more than 100 peer-reviewed journal articles and author of numerous reviews and book chapters on sarcomas. He co-authored the WHO Classification of Tumors of Bone and Soft Tissue Tumors (sarcomas) (2002 and 2012). He also has played key roles in the identification of therapeutic targets in gastrointestinal stromal tumor and dermatofibrosarcoma protuberans resulting in FDA approval of imatinib mesylate. His lab is currently focused on identifying therapeutic targets in epithelioid hemangioendothelioma. Contact Dr. Rubin at 216.445.5551 or at rubin2@ccf.org.
Alumni Connect

Dear Alumnus,

This is the fourth edition of our commitment to provide you with regular alumni communication. In this issue we bring you an update on L Building renovations.

L Building renovations completed

The Robert J. Tomsich Pathology & Laboratory Medicine Institute (RT-PLMI) is pleased to showcase our newly remodeled areas in the L Building. Hematopathology is now located in larger space on the third floor, with three sign-out rooms, a larger resident area, new laboratories, a conference room and remodeled offices. The previous Heme Path space in L1 now houses the Center for Pathology Education. The Center has a new residents’ and fellows’ headquarters, the MT and Cytotechnologist schools with a Wet Lab, a library and a new conference room with adjoining huddle rooms. The former residents’ headquarters in L2 became our new E-pathology area that services digital imaging for education and consults both regionally and worldwide. Automated chemistry and hematology and central processing in L1 are entirely remodeled.

We invite you to stop by to tour the newly remodeled L Building. We would be pleased to show you around.

We want to hear from you

Please send us your news and accomplishments to be featured in this “Alumni Connect” section in future issues of Pathology Innovations. If you prefer to receive an electronic version, please let us know by providing your preferred email address to ClientServices@ccf.org.

Fadi W. Abdul-Karim, MD, MMEd Vice-Chair, RT-PLMI Center for Pathology Education

Jonathan L. Myles, MD, Pathology and Laboratory Medicine Specialty Director

Cleveland Clinic Alumni Association Alumni Connect Steering Committee: Drs. Abdul-Karim and Myles, Daniel Kelly, Kathy Leonhardt, Emily Lopick, Paul Suchy, PhD, and Karl Theil, MD.
RT-PLMI well represented at USCAP

The 2015 USCAP annual meeting, the largest gathering of pathologists in the world, was held in Boston March 21–27. As has been true for many years, the presence of residents, fellows and faculty members from the Robert Tomsich Pathology and Laboratory Medicine Institute was strongly felt.

“We are proud of our residents, fellows and faculty members who contributed to the success of this highly prestigious meeting attended by more than 5,000 pathologists worldwide,” says John R. Goldblum, MD, Chair of the Department of Pathology.

Sixty-nine abstracts were presented from Cleveland Clinic residents, fellows and faculty. In the majority of these abstracts, RT-PLMI residents and fellows served as the first author for these presentations.

In addition to the strong presence in the poster and platform sessions, no other institution was as strongly represented in the meeting’s educational activities. Ten RT-PLMI faculty served as short course directors, which is one of the primary educational forums at the meeting. Three faculty members served as presenters or moderators for the evening subspecialty conferences, and five faculty members served as either moderators or presenters at the various companion society meetings held at the front end of the meeting. A number of faculty members served as scientific platform moderators, and two faculty served as course directors and/or presenters of USCAP special courses, including Carol Farver, MD, who served as course director and presenter for the residents’ workshop “Leadership, Collaboration and Change in Healthcare - Essential Skills,” and Tarik Elsheikh, MD, who served as course director and presenter for the special course, “Basic Principles in Cytology.” John Goldblum, MD, served as the moderator for the second annual “Hot Topics in Gastrointestinal Pathology” luncheon meeting.

Several individuals also hold key positions within the USCAP or related companion societies. This past year, Dr. Elsheikh served as chair of the USCAP Foundation Committee. Brian Rubin, MD, PhD, and Deborah Chute, MD, are members of the Vogel Award and Stowell-Orbison Award Committees, respectively, and Rish Pai, MD, PhD, and Drs. Elsheikh and Goldblum serve as members of the Innovative Educational Products Development Committee. The Education Committee is the “lifeblood” of the USCAP, and Cleveland Clinic has a major presence on this committee, including Jesse McKenney, MD, who served as the outgoing short course coordinator, Steven Billings, MD, who will be serving as the incoming short course coordinator, and Dr. Farver, who is the newest member of the Education Committee. Dr. Billings is also an important member of the Education Committee’s Unique Live Course Offerings Subcommittee. In addition, Dr. Goldblum served as the USCAP’s President Elect and member of the Board of Directors.

Sara Falzarano, MD (left photo), a pathology resident, and Angela Collie, MD, PhD (right photo), who recently completed her dermatopathology fellowship, were among the many Cleveland Clinic residents, faculty and staff members who represented the Robert Tomsich Pathology and Laboratory Medicine Institute in 69 abstracts presented at the USCAP meeting in Boston.
Pathology services integrated with regional hospitals

Earlier this year the Robert J. Tomsich Pathology & Laboratory Medicine Institute began performing diagnosis of all biopises and surgical/cytology specimens for Cleveland Clinic patients – no matter their location – by pathologists with subspecialized expertise, including gynecologic pathology.

In the last few years, regional hospital pathology technical services have been successfully consolidated to Cleveland Clinic’s Hillcrest and Fairview hospitals. To complete this progression and move to full subspecialty pathology practice, the technical pathology and gynecologic cytology processing services performed at Hillcrest and Fairview will be moved to main campus.

Dr. Elsheikh named President of PSC

Tarik Elsheikh, MD, was recently elected to a two-year term as President of the Papanicolaou Society of Pathology (PSC), an international physician association dedicated to education in cytopathology and small biopsy histology. Society members actively practice cytopathology and surgical pathology at prestigious institutions throughout the United States and worldwide.

RT-PLMI News

Ana Bennett, MD, is the new Director of Pathology Operations.

Mark Melargano, MD, is the new Director of Regional Operations.

Leal Herlitz, MD, is the new Director of Medical Kidney Pathology.

Melissa Piliang, MD, joins Steven Billings, MD, as Co-section Head of Dermatopathology.

The June 1 issue of The Dark Report featured an article, “Lab Test Utilization Delivers Big Gains at Cleveland Clinic,” with Gary Procop, MD.