A 2018 Reference Guide to the Banff Classification of Renal Allograft Pathology

Candice Roufossé, MD, PhD,1,2 Naomi Simmonds, MD,3 Marian Claesen-van Groningen, MD, PhD,4 Mark Haas, MD, PhD,5 Kammi J. Henriksen, MD,6 Catherine Horsfield, MD,5 Alexandre Loupy, MD,7 Michael Mengel, MD,8 Agnieszka Perkowska-Ptasinska, MD,9 Marion Rabant, MD, PhD,10 Lorraine C. Racusen, MD,11 Kim Solze, MD,8 and Jan U. Becker, MD12

Abstract: The Banff Classification of Allograft Pathology is an international consensus classification for the reporting of biopsies from solid organ transplants. Since its initial conception in 1991 for renal transplants, it has undergone review every 2 years, with attendant updated publications. The rapid expansion of knowledge in the field has led to numerous revisions of the classification. The resultant dispersal of relevant content makes it difficult for novices and experienced pathologists to faithfully apply the classification in routine diagnostic work and in clinical trials. This review shall provide a complete and simple illustrated reference guide of the Banff Classification of Kidney Allograft Pathology based on all publications including the 2017 update. It is intended as a concise desktop reference for pathologists and clinicians, providing definitions, Banff Lesion Scores and Banff Diagnostic Categories. An online website reference guide hosted by the Banff Foundation for Allograft Pathology (www.banfffoundation.org) is being developed, which will be updated with future refinement of the Banff Classification from 2019 onward.

Since its first consensus meeting in 1991,1 the Banff Classification of Allograft Pathology has provided a framework for the reporting of renal allograft biopsies. It was the first classification system of its kind and answered the need for an international consensus on renal transplant biopsy reporting, providing guidance for clinical diagnosis and enabling meaningful comparison between research studies and clinical trials investigating the diagnosis, treatment and outcome in kidney transplantation. The Banff Classification has since been further strengthened by evidence-informed biannual updates elaborated during open international expert meetings.2 As a result, the Banff Classification of Allograft Pathology has become the predominant classification system used worldwide.3

A total of 14 meetings reported in 10 articles reflect the developments of the Banff Classification from the first consensus meeting in 1991 to the recently published consensus

Received 19 February 2018. Revision received 7 May 2018.
Accepted 23 May 2018.
1 Department of Medicine, Imperial College, London, United Kingdom.
2 North West London Pathology, London, United Kingdom.
3 Department of Histopathology, Guy’s and St. Thomas’ National Health Service Foundation Trust, London, United Kingdom.
4 Department of Pathology, Erasmus MC, Rotterdam, The Netherlands.
5 Department of Pathology, Cedars-Sinai Medical Center, Los Angeles, CA.
6 Department of Pathology, University of Chicago, Chicago, IL.
7 Paris Translational Research Center for Organ Transplantation, Paris, France.
8 Department of Laboratory Medicine and Pathology, University of Alberta, Edmonton, Canada.
9 Department of Transplantology, Nephrology and Internal Diseases, Medical University of Warsaw, Warsaw, Poland.
10 Department of Pathology, Necker Hospital University Paris Descartes, Paris, France.
11 The Johns Hopkins University, Baltimore, MD.
12 Institute of Pathology, University Hospital of Cologne, Cologne, Germany.

C.R.’s and N.S.’s contribution to this research is supported by the National Institute for Health Research (NIHR) Biomedical Research Centre based at Imperial College Healthcare NHS Trust and Imperial College London. A.L. is supported by a research grant ATIP AVENIR from the National French Institute of Research. J.U.B. is supported by the European Rare Kidney Disease Network (ERKNet) and by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation - BE-3801).
The authors declare no conflicts of interest.
All authors contributed the discussion of the content, the collation of images and illustrations and to writing of the article.
Correspondence: Jan U. Becker, MD, Institute of Pathology, University Hospital of Cologne, Kerpener Str. 62, 50937, Germany. (janbecker@gmx.com).
Supplemental digital content (SDC) is available for this article. Direct URL citations appear in the printed text, and links to the digital files are provided in the HTML text of this article on the journal’s Web site (www.transplantjournal.com).
Copyright © 2018 The Author(s). Published by Wolters Kluwer Health, Inc. This is an open access article distributed under the Creative Commons Attribution License 4.0 (CCBY), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.
ISSN: 0041-1337/18/10211-1795
DOI: 10.1097/TP.0000000000002366
after the 2017 meeting in Barcelona, Spain. Each of these iterations provides a short summary of the meeting and contributes to the classification in a cumulative fashion. The dispersal of both relevant and outdated content over 10 articles could make access to the Banff Classification difficult for beginners and experts and has created ambiguities in the past. Yet, accessibility and clarity are of utmost importance not only for clinical practice and research but also for the Banff Classification itself to evolve through accountability, critique, and change. To improve on these aspects, the Rules and Dissemination Banff Working Group was initiated at the last Banff meeting held in Barcelona, Spain in March 2017. With a scope beyond the helpful syllabus provided by the Banff group in the online supplement of the 2015 update and incorporating the latest changes introduced in the 2017 update, the aim of this Working Group is to collate all current content of the Banff Classification and improve its accessibility. A systematic inventory of the content is given in Figure 1. This practical guide is based on all content up to the 2017 update as the first output of our Working Group. It is divided in the following sections: a brief guide about the histopathological and serological work-up; a list of Banff Lesion Scores (previously known as components, eg, Banff f for tubulitis) with their current definitions, practical tips for their application and illustrative figures (see definitions below and thresholds in Table 2); and a list of Banff Diagnostic Categories in Table 1. Moreover, we provide a list of Additional Diagnostic Parameters, which need to be considered in addition to Banff Lesion Scores to reach a Banff Diagnostic Category (Table 3). Examples for these include “Severe Peritubular Capillary Basement Membrane Multilayering” which is among the criteria for antibody-mediated rejection (AMR) chronicity. A glossary of terms is provided as Supplemental Digital Content (see Glossary of Terms, SDC, http://links.lww.com/TP/B604), explaining important concepts and terminology underlying the Banff Classification. Lastly, we provide a critical appraisal of areas of the Banff Classification that require clarification and provide an outlook for future developments. All terms from the Banff Classification will be given in capitals for clarity, all abbreviations for Banff lesion scores will be given in italic typeface.

We hope this Banff 101 will serve as a handy reference for the clinicians and the pathologists, until the entire updated content appears online with the 2019 update of the Banff Classification of Renal Allograft Pathology, replacing this guide.

**DIAGNOSTIC WORK-UP OF BIOPSIES**

A kidney transplant biopsy should fulfill the criteria for specimen adequacy (see Glossary of Terms, SDC, http://links.lww.com/TP/B604) detailed in the Banff 1997 update. C4d staining is considered indispensable, either as immunofluorescence (IF) on fresh frozen or immunohistochemistry (IHC) on paraffin-embedded tissue. The paraffin block should be cut in several numbered level sections examined with hematoxylin-eosin, periodic acid-Schiff (PAS), trichrome-elastic and Jones or methenamine silver stains. Immunohistochemistry staining for simian virus-40, cross-reacting with BK virus is highly recommended when indicated. Where available, minute portions of cortex should be embedded for transmission electron microscopy (EM).

Depending on clinical and histopathological findings a complete nephropathological work-up including staining for immunoglobulin heavy and light chains and complement split products might be necessary to rule out or confirm a diagnosis of glomerulonephritis. Other ancillary staining might be necessary as for native kidney biopsies to establish specific recurrent or de novo kidney diseases (eg, Congo red stain). Serological testing for donor-specific antibodies (DSAs) should be performed as described in respective consensus documents. Ancillary molecular tests, based on tissue and body fluids, are emerging.

Preimplantation biopsies should be obtained, processed, and reported as described by the Banff Working Group on Preimplantation Biopsies.

**BANFF LESION SCORES**

Banff Lesion Scores assess the presence and the degree of histopathological changes in the different compartments of renal transplant biopsies, focusing primarily but not exclusively on the diagnostic features seen in rejection. These Banff Lesion Scores are not by themselves sufficient to reach the various Banff Diagnostic Categories in Table 1; the Additional Diagnostic Parameters—histopathological, molecular, serological and/or clinical—may be required to determine the diagnosis. For each Banff Lesion Score we give the current consensus definitions below. As new knowledge emerges, these might be refined for the forthcoming Banff 2019 update. A synopsis of their semiquantitative thresholds is given in Table 2. However, use of this threshold table without knowledge of the precise definitions and regulatory statutes underlying each Banff Lesion Score is strongly discouraged.

**Banff Lesion Score i (Interstitial Inflammation)**

This score evaluates the degree of inflammation in non-scarrred areas of cortex, which is often a marker of Acute T Cell–Mediated Rejection (TCMR). As per the Banff update from 1997, areas that must not be considered for Banff Lesion Score i are “fibrotic areas, the immediate subcapsular cortex, and the adventitia around large veins and lymphatics.” As can indirectly be derived from the definition of Banff Lesion Score ti in the 2007 update of the Banff classification, nodular infiltrates, in unscarred cortex, are also considered for Banff Lesion Score i. An asterisk shall be
**TABLE 1.**

Banff Diagnostic Categories form the core of the Banff Classification of Renal Allograft Pathology

**Category 1: Normal Biopsy Or Nonspecific Changes**

Requires exclusion of any diagnosis from the Banff Diagnostic Categories 2-4, 6 below.

**Category 2: Antibody-mediated changes**

Use the Diagnostic Criteria Groups (right column) to reach 1 Diagnosis (left column)

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Criteria Group 1 AMR activity:</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4d Staining Without Evidence of Rejection</td>
<td>- Banff Lesion Score g &gt; 0 in the absence of glomerulonephritis and/or Banff Lesion Score ptc &gt; 0 in the absence of TCMR or Borderline</td>
</tr>
<tr>
<td>Banff Lesion Score C4d &gt; 1 (IF on fresh frozen tissue) OR C4d &gt; 0 (IHC on paraffin-embedded tissue) AND Banff Lesion Scores t0, v0, no arterial intimal fibrosis with mononuclear cell inflammation in fibrosis and formation of neointima, no criterion from group 1 (AMR activity), no criterion from group 4 (histologic features of AMR chronicity), no increased expression of thoroughly validated gene transcripts/classifiers in the biopsy tissue strongly associated with AMR</td>
<td>- Banff Lesion Score v &gt; 0</td>
</tr>
<tr>
<td>- Acute Thrombotic Microangiopathy In The Absence Of Any Other Cause (Figure 18)</td>
<td></td>
</tr>
<tr>
<td>- Active T cell-mediated rejection (TCMR; Diagnostic Category 4). If Borderline, acute TCMR, or infection are present, Banff Lesion Scores (g + ptc) &gt; 1 is not sufficient and Banff Lesion Score g &gt; 1 is required.</td>
<td></td>
</tr>
<tr>
<td>- Increased Expression Of Thoroughly Validated Gene Transcripts/Classifiers In The Biopsy Tissue Strongly Associated With AMR</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Criteria Group 2 Antibody interaction with tissue:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active AMR</td>
<td>- Banff Lesion Score C4d &gt; 1 (IF on fresh frozen tissue) or C4d &gt; 0 (IHC on paraffin-embedded tissue)</td>
</tr>
<tr>
<td>No criterion of AMR chronicity (Criteria Group 4) AND At least 1 criterion from Criteria Group 1 (AMR activity) AND At least 1 criterion from Criteria Group 2 (antibody interaction with tissue) AND At least 1 criterion from Criteria Group 3 (DSA or equivalents)</td>
<td>- At least moderate MVI (g + ptc &gt; 1) in the absence of recurrent or de novo glomerulonephritis; Borderline (Diagnostic Category 3) or acute T cell-mediated rejection (TCMR; Diagnostic Category 4). If Borderline, acute TCMR, or infection are present, Banff Lesion Scores (g + ptc) &gt; 1 is not sufficient and Banff Lesion Score g &gt; 1 is required.</td>
</tr>
<tr>
<td>- Increased Expression Of Thoroughly Validated Gene Transcripts/Classifiers In The Biopsy Tissue Strongly Associated With AMR</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Criteria Group 3 DSA or equivalents:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chronic Active AMR</td>
<td>- DSA (anti-HLA or other specificity)</td>
</tr>
<tr>
<td>At least 1 feature of AMR chronicity (Criteria Group 4) AND At least 1 criterion of antibody interaction with tissue (Criteria Group 2) AND At least 1 criterion of DSA or equivalents (Criteria Group 3)</td>
<td>- Banff Lesion Score C4d &gt; 1 (IF on fresh frozen tissue) or C4d &gt; 0 (IHC on paraffin-embedded tissue)</td>
</tr>
<tr>
<td>- Increased Expression Of Thoroughly Validated Gene Transcripts/Classifiers In The Biopsy Tissue Strongly Associated With AMR</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Criteria Group 4 Histologic features of AMR chronicity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chronic AMR</td>
<td>- Banff Lesion Score cg &gt; 0 (by LM or EM, if available), excluding biopsies with evidence of chronic thrombotic microangiopathy</td>
</tr>
<tr>
<td>Banff 2017 permits the use of this term for biopsy specimens showing TG and/or peritubular capillary basement membrane multilayering in the absence of criterion of current/recent antibody interaction with the endothelium (Criteria Group 2) but with a prior documented diagnosis of Active or Chronic Active AMR or documented prior evidence of DSA</td>
<td>- 7 or more layers in 1 cortical peritubular capillary and 5 or more in 2 additional capillaries, avoiding portions cut tangentially by EM, if available (Severe Peritubular Capillary Basement Membrane Multilayering, see Figure 19)</td>
</tr>
<tr>
<td>- Arterial Intimal Fibrosis Of New Onset, Excluding Other Causes; Leukocytes Within The Sclerotic Intima Favor Chronic AMR If There Is No Prior History Of Biopsy-Proven TCMR but are not required</td>
<td></td>
</tr>
</tbody>
</table>

Continued next page
TABLE 1. (Continued)

**Category 3: Suspicious (Borderline) For Acute TCMR**

- Foci of Banff Lesion Score $t > 0$ AND Banff Lesions Score $i \leq 1$ (retaining the Banff Lesion Score $i^1$ threshold from Banff 2005 is permitted but it must be made transparent in the methods section of reports and publications)
- OR
- Foci of Banff Lesion Score $t^1$ AND Banff Lesion Score $i \geq 2$

**Category 4: TCMR**

- Acute TCMR IA
  - Banff Lesion Score $i \geq 2$
  - AND
  - Banff Lesion Score $i^2$
- Acute TCMR IB
  - Banff Lesion Score $i \geq 2$
  - AND
  - Banff Lesion Score $i^3$
- Acute TCMR IIA
  - Banff Lesion Score $v^1$ regardless of Banff Lesion Scores $i$ or $t$
- Acute TCMR IIB
  - Banff Lesion Score $v^2$ regardless of Banff Lesion Scores $i$ or $t$
- Acute TCMR III
  - Banff Lesion Score $v^3$ regardless of Banff Lesion Scores $i$ or $t$

**Chronic Active TCMR Grade IA**

- Banff Lesion Score $t^1_i \geq 2$
  - AND
  - Banff Lesion Score $i-IFTA \geq 2$, other known causes of $i$-IFTA (eg, pyelonephritis, BK-virus nephritis etc.) ruled out
  - AND
  - Banff Lesion Score $t^2_i$
- Chronic Active TCMR Grade IB
  - Banff Lesion Score $t^1_i \geq 2$
  - AND
  - Banff Lesion Score $i-IFTA \geq 2$, other known causes of $i$-IFTA ruled out
  - AND
  - Banff Lesion Score $t^3_i$

**Category 5: IFTA**

**Grade I (Mild)**

- Banff Lesion Score $c^1_i$
- OR
- Banff Lesion Score $c^1_t$
added to Banff Lesion Score \( i \) (eg, \( i_1^* \)), “if there are more than 5% to 10% of eosinophils, neutrophils or plasma cells”. Exemplary lesions are shown in Figure 2.

\( i_0 \)—No inflammation or in less than 10% of unscarred cortical parenchyma.

\( i_1 \)—Inflammation in 10 to 25% of unscarred cortical parenchyma.

\( i_2 \)—Inflammation in 26 to 50% of unscarred cortical parenchyma.

\( i_3 \)—Inflammation in more than 50% of unscarred cortical parenchyma.\(^{11}\)

**Banff Lesion Score \( t \) (Tubulitis)**

This Banff Lesion Score evaluates the degree of inflammation within the epithelium of the cortical tubules. As per the Banff 2003 update “Tubulitis—the presence of mononuclear cells in the basolateral aspect of the renal tubule epithelium” is one of the defining lesion of TCMR in kidney transplants.\(^{6}\) According to Banff 1997, in tubules cut longitudinally, the score shall be determined as the number of mononuclear cells per 10 tubular epithelial cells, which is the average number of epithelial cells per tubular cross-section (Figure 3). Tubulitis must be present in at least 2 foci. We have emphasized this by rephrasing the criteria for Banff Lesion Score \( t_0 \) below; the most severely affected tubule determines the score.\(^{3,11}\) Please note also that we have returned from the altered definition with “leukocytes” in the Banff 2015 update\(^{11}\) to “mononuclear cells” as given in the 1997 update.\(^{5}\)

According to the most recent Banff update from 2017, for Acute TCMR Grade IA, IB and Chronic Active TCMR Grade IA and IB but not Borderline (Banff Diagnostic Category 3), tubulitis is considered in all but severely atrophic cortical tubules. Tubulitis in severely atrophic tubules does not count toward a diagnosis of either Borderline, Acute or Chronic Active TCMR, and severely atrophic tubules are defined by a diameter of less than 25% of that of unaffected or minimally affected tubules on the biopsy, often with an undifferentiated appearing, cuboidal or flattened epithelium (or in some cases even loss of epithelium with denudation of the tubular basement membrane), and pronounced wrinkling and/or thickening of the tubular basement membrane. This definition of severely atrophic tubules also includes very small, endocrine-like tubules with very narrow lumens, although the basement membranes of the latter may not be thickened.\(^{12}\) An example of tubulitis in various stages of tubular atrophy is shown in Figure 4.

\( t_0 \)—No mononuclear cells in tubules or single focus of tubulitis only.

\( t_1 \)—Foci with 1 to 4 mononuclear cells/tubular cross section (or 10 tubular cells).

\( t_2 \)—Foci with 5 to 10 mononuclear cells/tubular cross section (or 10 tubular cells).

\( t_3 \)—Foci with \( >10 \) mononuclear cells/tubular cross section or the presence of \( \geq 2 \) areas of tubular basement membrane destruction accompanied by \( i_2/i_3 \) inflammation and \( t_2 \) elsewhere.\(^{12}\)

**Banff Lesion Score \( v \) (Intimal Arteritis)**

This Banff Lesion Score evaluates the presence and the degree of inflammation within the arterial intima. Arteries are defined as having at least 2 layers of smooth muscle cells in the media (Glossary of Terms, SDC, http://links.lww.com/TP/B604). Note that intimal arteritis (also referred to as
endothelialitis and endarteritis) is defined by the presence of inflammatory cells, mainly lymphocytes and monocytes, in the subendothelial space of 1 or more arteries. One such cell suffices. Examples of this lesion are shown in Figure 5.

Intimal arteritis is a feature seen in both Acute TCMR and Active AMR. For Banff Lesion Score $v$, the most severely affected artery dictates the score. Similar lesions in arterioles are only coded as an asterisk behind the Banff Lesion Score and are disregarded for Banff Lesion Score $v$. Infiltrates buried deeper in the intima are not considered for the $v$ Banff Lesion Score but have been recognized as Chronic Active TCMR since the 2005 update, and graded in the 2017 update as Grade II. In the presence of tubulointerstitial hemorrhage (see Glossary of Terms, SDC, http://links.lww.com/TP/B604) and/or infarct (see Glossary of Terms, SDC, http://links.lww.com/TP/B604) an asterisk “*” is attached to the Banff Lesion Score $v$ (eg, Banff $v0^*, v2^*$).

$v0$—No arteritis.

$v1$—Mild to moderate intimal arteritis in at least 1 arterial cross section.

$v2$—Severe intimal arteritis with at least 25% luminal area lost in at least 1 arterial cross section.

### TABLE 2. This is a synopsis of the thresholds for all Banff Lesion Scores

<table>
<thead>
<tr>
<th>Banff lesion score, Abbreviation</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interstitial inflammation $i$</td>
<td>$&lt;$10%</td>
<td>10-25%</td>
<td>26-50%</td>
<td>$&gt;$50</td>
</tr>
<tr>
<td>Tubulitis $t$</td>
<td>None</td>
<td>1-4/tubular cross section or 10 tubular epithelial cells</td>
<td>5-10</td>
<td>$&gt;$10 or foci of tubular basement membrane destruction with $i \geq 2$ and $t2$ elsewhere Transmural and/or fibrinoid change and medial smooth muscle necrosis</td>
</tr>
<tr>
<td>Intimal arteritis $v$</td>
<td>None</td>
<td>$&lt;$25% luminal area lost</td>
<td>$\geq$25% luminal area lost</td>
<td></td>
</tr>
<tr>
<td>Glomerulitis $g$</td>
<td>None</td>
<td>$&lt;$10% of</td>
<td>$\geq$10% of</td>
<td>$\geq$25% luminal area lost</td>
</tr>
<tr>
<td>Peritubular capillaritis $ptc$</td>
<td>$&lt;$3 leukocytes/PTC</td>
<td>$\geq$1 leukocyte in $\geq$10% of PTCs with max. of 3-4/PTC</td>
<td>$\geq$1 leukocyte in $\geq$10% of PTCs with max. of 5-10/PTC</td>
<td>$\geq$1 leukocyte in $\geq$10% of PTCs with max. of $&gt;$10/PTC</td>
</tr>
<tr>
<td>C4d $C4d$</td>
<td>None</td>
<td>$&lt;$10%</td>
<td>10-50%</td>
<td>$&gt;$50</td>
</tr>
<tr>
<td>Interstitial fibrosis $ci$</td>
<td>$\leq$5%</td>
<td>6-25%</td>
<td>26-50%</td>
<td>$&gt;$50</td>
</tr>
<tr>
<td>Tubular atrophy $ct$</td>
<td>$\leq$25%</td>
<td>26-50%</td>
<td>$&gt;$50</td>
<td></td>
</tr>
<tr>
<td>Vascular fibrous intimal thickening $cv$</td>
<td>None</td>
<td>$\leq$25%</td>
<td>$&gt;$50</td>
<td></td>
</tr>
<tr>
<td>GBM double contours $cg$</td>
<td>None</td>
<td>$1a$: only by EM</td>
<td>$1b$: $\leq$25% by LM</td>
<td>$&gt;$50</td>
</tr>
<tr>
<td>Mesangial matrix expansion $mm$</td>
<td>None</td>
<td>$\leq$25%</td>
<td>26-50%</td>
<td>$&gt;$50</td>
</tr>
<tr>
<td>Arteriolar hyalinosis $ah$</td>
<td>None</td>
<td>Mild to moderate in $\geq$1</td>
<td>Moderate to severe in $&gt;1$</td>
<td>Severe in many circumferential</td>
</tr>
<tr>
<td>Hyaline arteriolar thickening $aah$</td>
<td>None</td>
<td>$1$ without circumferential</td>
<td>$\geq$1 without circumferential</td>
<td>$&gt;$50</td>
</tr>
<tr>
<td>Total inflammation $ti$</td>
<td>$&lt;$10%</td>
<td>10-25%</td>
<td>26-50%</td>
<td>$&gt;$50</td>
</tr>
<tr>
<td>Inflammation in the area of IFTA $i$-IFTA</td>
<td>$&lt;10%$</td>
<td>10-25%</td>
<td>26-50%</td>
<td>$&gt;$50</td>
</tr>
</tbody>
</table>

The user of this table should be familiar with the exact definitions underlying each individual Banff Lesion Score. Reliance on these thresholds alone without consideration of the regulatory statutes behind these scores is strongly discouraged. max.: maximum; PTC, peritubular capillary.

![FIGURE 2. Banff Lesion Score i (Interstitial Inflammation in nonscarred areas of the cortex). A, Interstitial inflammation in nonscarred areas of the cortex. This Banff Lesion Score, often a marker of TCMR, ranges from 0 to 3, based on the percentage of nonscarred cortex involved, and is usually dominated by mononuclear cells in the case of Acute TCMR. Note the contrast between the noninfiltrated interstitium in the right half of the micrograph and the infiltrate in the edema between the tubules on the left (long arrow). PAS, original magnification ×400. B, An example of plasma cell rich interstitial inflammation. If the infiltrate comprises more than 5% to 10% of either eosinophils, neutrophils or plasma cells an asterisk is added to the Banff Lesion Score i (eg, i*). H&E, hematoxylin and eosin, original magnification ×400.](www.transplantjournal.com)
v3—Transmural arteritis and/or arterial fibrinoid change and medial smooth muscle necrosis with lymphocytic infiltrate in vessel.11

**Banff Lesion Score g (Glomerulitis)**

This Banff Lesion Score evaluates the degree of inflammation within glomeruli (Figure 6). Glomerulitis is a form of microvascular inflammation (MVI) and is a feature of activity and antibody interaction with tissue in AMR. It can also be seen in recurrent or de novo glomerulonephritis which must be excluded by appropriate immunostains and EM.

Banff Lesion Score g is determined by the proportion of glomeruli showing glomerulitis defined as "complete or partial occlusion of 1 or more glomerular capillary by leukocyte infiltration and endothelial cell enlargement." Leukocytes include polymorphonuclear cells and mononuclear cells. Both endothelial cell enlargement and leukocyte(s) must contribute to the complete or partial occlusion. The denominator in this proportion is the number of nonsclerosed glomeruli in the biopsy.

- g0—No glomerulitis.
- g1—Segmental or global glomerulitis in less than 25% of glomeruli.
- g2—Segmental or global glomerulitis in 25 to 75% of glomeruli.
- g3—Segmental or global glomerulitis in more than 75% of glomeruli.11

**Banff Lesion Score ptc (Peritubular Capillaritis)**

This Banff Lesion Score evaluates the degree of inflammation within peritubular capillaries (PTCs). Together with glomerulitis, peritubular capillaritis constitutes MVI as a feature of Active AMR or Chronic Active AMR. Peritubular capillaritis can be observed with pure Acute TCMR or Borderline as well.

According to the Banff 2005 update, the Banff Lesion Score ptc is determined by the most severely involved PTC (Figure 7). Peritubular capillaries are by definition found in the cortex, their medullary equivalent are medullary vasa recta. The number of luminal inflammatory cells includes polymorphonuclear and mononuclear leukocytes, with an asterisk "*" used to indicate only mononuclear cells and absence of neutrophils. The extent of the PTC inflammation in the biopsy should be documented, either as focal (10-50% of cortical area) or diffuse (>50% of cortical area), but this does not contribute to the score. The presence of associated PTC dilatation may also be noted. Areas affected by acute

---

**FIGURE 3.** Banff Lesion Score t (tubulitis) in nonatrophic or mildly atrophic tubules. These images display various degrees of tubulitis which is characterized by the presence of mononuclear cells on the basolateral aspect of the tubular epithelial cells, within the confines of the basement membrane. Mononuclear cells (long and short arrows) are noticeable by their characteristic halo and smaller nucleus and more condensed chromatin compared to tubular epithelial cells. A, Banff Lesion Score t0—Cortical tubules without tubulitis which would be scored as t0. H&E, original magnification ×200. B, Banff Lesion Score t1—defined as foci of 1-4 mononuclear cells (arrows) per tubular cross section or per 10 tubular epithelial cells. PAS, original magnification ×400. C, Banff Lesion Score t2—defined as 5 to 10 mononuclear cells per tubular cross section or per 10 epithelial cells (long arrows). Note that the tubule to the left displays mild tubulitis (short arrows), but the most severely affected tubule dictates the score. PAS, original magnification ×400. D, Banff Lesion Score t3—defined as foci with >10 mononuclear cells/tubular cross section. Note that for this particular tubule the denominator is per 10 tubular epithelial cells as this tubule is sectioned longitudinally. PAS, original magnification ×400.
nosed as Acute TCMR Grade IB. PAS, original magnification atrophic. The mononuclear tubulitis in this particular tubule must be the tubule with the diameter of 20 μm marked in red is heavily infiltrated with mononuclear cells (gray arrows). In contrast, 50% of 48ullary capillaries (stains (PAS, silver). Inflammatory cells within veins and medullary specific inflammation should not be scored. Inflammatory pyelonephritis or necrosis and subcapsular cortex with non-affected tubules with their diameter marked in black. Their mean diameter in this image would be around 48 μm. The tubule with the diameter marked in gray has a diameter of 27 μm which is more than 50% of 48 μm. Thus, this tubule would still qualify as mildly atrophic. It is heavily infiltrated with mononuclear cells (gray arrows). In contrast, the tubule with the diameter of 20 μm marked in red is moderately atrophic. The mononuclear tubulitis in this particular tubule must be scored toward Banff Lesion Score t in this biopsy which was diagnosed as Acute TCMR Grade IB. PAS, original magnification ×400.

Pyelonephritis or necrosis and subcapsular cortex with non-specific inflammation should not be scored. Inflammatory cells within PTCs must be distinguished from interstitial inflammation by careful examination of basement membrane stains (PAS, silver). Inflammatory cells within veins and medullary capillaries (vasa recta) should not be scored. Consequently, peritubular capillaritis and Banff Lesion Score ptc can only be assessed in the cortex after exclusion of areas of pyelonephritis and infarcted areas and exclusion of areas close to lymphoid aggregates to avoid confusion with lymphatic vessels. Banff Lesion Score ptc should not be based on longitudinally cut PTCs. Peritubular capillaries in areas affected by tubular atrophy and interstitial fibrosis must explicitly be considered for this Banff Lesion Score. Note that we have simplified the definition of ptc0 from the original version in the Banff 2017 update.12

ptc0—Maximum number of leukocytes <3.
ptc1—At least 1 leukocyte cell in ≥10% of cortical PTCs with 3-4 leukocytes in most severely involved PTC.
ptc2—At least 1 leukocyte in ≥10% of cortical PTC with 5-10 leukocytes in most severely involved PTC.
ptc3—At least 1 leukocyte in ≥10% of cortical PTC with >10 leukocytes in most severely involved PTC.11

Banff Lesion Score C4d

This score evaluates the extent of staining for C4d on endothelial cells of PTCs and medullary vasa recta by IF on snap frozen sections of fresh tissue or IHC on formalin-fixed and paraffin-embedded tissue. Although Banff 2007 states that areas of tubular atrophy and interstitial fibrosis have reduced PTC density that could affect the extent of staining,15 scoring of C4d in such cortical areas is not excluded.3 Scoring of C4d staining is based on the percentage of peritubular capillaries and vasa recta that has a linear, circumferential staining pattern (Figure 8). The minimal sample for evaluation is 5 high-power fields of cortex and/or medulla without scarring or infarction. C4d must not be scored in areas of infarction. On IF, staining should be at least 1+ in intensity.8 Strong staining is not required for a positive reading for IHC.11 In Banff Lesion Score this lesion score evaluates the extent of cortical fibrosis. The Banff Classification has never given a precise definition for individual areas of interstitial fibrosis (Figure 9). The reason for this is that Banff Lesion Score ci was meant to purely reflect the cortex composed of fibrous tissue, which does not necessarily correspond to areas that a pathologist would pick up as a patch of pathological tubulointerstitial fibrosis. The fraction of fibrous tissue in the cortex was considered as up to 5% for normal kidneys, hence the difference in cut-offs between ci1 and ci1. A Working Group on this topic has produced useful reference guides (Figures 10 and 11).17

ci0—Interstitial fibrosis in up to 5% of cortical area.

ci1—Interstitial fibrosis in 6 to 25% of cortical area (mild interstitial fibrosis).

ci2—Interstitial fibrosis in 26 to 50% of cortical area (moderate interstitial fibrosis).

ci3—Interstitial fibrosis in >50% of cortical area (severe interstitial fibrosis).11

Banff Lesion Score ct (Tubular Atrophy)

This Banff Lesion Score evaluates the extent of cortical tubular atrophy which is usually tightly associated with the areas affected with interstitial fibrosis (Figure 9). Both correlate with time posttransplantation in the setting of progressive disease of any cause. Accordingly, neither Banff Lesion Scores ct nor ci have diagnostic specificity, but both have significant correlation with allograft function and prognosis. Historically, the Banff classification has defined tubular atrophy as reflected in the Banff Lesion Score ct in the 1995 update as tubules with a thickened basement membrane or a reduction of greater than 50% in tubular diameter. Banff Lesion Score ct is still based on this definition of tubular atrophy. The definitions of moderate and severe atrophy from the Banff 2017 update are irrelevant for Banff Lesion Score ct. In the following definition, we have omitted the designation as

FIGURE 4. Banff Lesion Score t (tubulitis) in moderately atrophic tubules. In biopsies with Banff Lesion Scores i, t, and -IFTA sufficient for a diagnosis of Acute TCMR Grade IA, IB or Chronic Active TCMR Grade IA and IB, Banff Lesion Score t must also be scored in moderately atrophic cortical tubules. Moderately atrophic tubules are defined as having less than 50% down to 25% of the diameter of the surrounding "unaffected or minimally affected [cortical] tubules in the biopsy".12 This example shows such unaffected or minimally affected tubules with their diameter marked in black. Their mean diameter in this image would be around 48 μm. The tubule with the diameter marked in gray has a diameter of 27 μm which is more than 50% of 48 μm. Thus, this tubule would still qualify as mildly atrophic. It is heavily infiltrated with mononuclear cells (gray arrows). In contrast, the tubule with the diameter of 20 μm marked in red is moderately atrophic. The mononuclear tubulitis in this particular tubule must be scored toward Banff Lesion Score t in this biopsy which was diagnosed as Acute TCMR Grade IB. PAS, original magnification ×400.
mild for ct1, moderate for ct2 and severe for ct3 which was still included in the Banff 2015 update to avoid confusion between the definition of atrophy for an individual tubule as described above and the extent of tubular atrophy reflected in the Banff Lesion Score ct.

c0—No tubular atrophy.
c1—Tubular atrophy involving up to 25% of the area of cortical tubules.
c2—Tubular atrophy involving 26 to 50% of the area of cortical tubules.
c3—Tubular atrophy involving in >50% of the area of cortical tubules.11

Banff Lesion Score cv (Vascular Fibrous Intimal Thickening)

This Banff Lesion Score reflects the extent of arterial intimal thickening in the most severely affected artery (see Definition of Terms, SDC, http://links.lww.com/TP/B604), not the average of all arteries.1 It does not discriminate between bland arterial intimal fibrosis and fibrosis containing leukocytes (Figure 12), although the latter is more likely to reflect chronic rejection (AMR and/or Chronic Active TCMR Grade II).12 A visual analog scale for application in daily practice is provided in Figure 13.

c0—No chronic vascular changes.
c1—Vascular narrowing of up to 25% luminal area by fibrointimal thickening.
c2—Vascular narrowing of 26 to 50% luminal area by fibrointimal thickening.
c3—Vascular narrowing of more than 50% luminal area by fibrointimal thickening.11

Banff cg Score (Glomerular Basement Membrane Double Contours)

Banff Lesion Score cg is based on the presence and extent of glomerular basement membrane (GBM) double contours or multilamination in the most severely affected glomerulus (Figure 14). Scoring should be carried out on PAS or silver stains; a designation as cg1a requires transmission EM to exclude cg0. With Banff Lesion Score cg > 0 (including both cg1a and cg1b), a diagnosis of transplant glomerulopathy (TG) (see Glossary of Terms, SDC, http://links.lww.com/TP/B604) can be made, if other causes can be excluded. Banff Lesion Score cg > 0 can be a feature of Chronic AMR or...
FIGURE 7. Banff Lesion Score ptc (peritubular capillaritis). Peritubular capillaritis is a form of MVI and a feature of AMR activity. Each image demonstrates the various ptc Scores which are in themselves determined by the number of inflammatory cells present within capillary lumina. A, Banff Lesion Score ptc1—Mild peritubular capillaritis defined as at least 1 cell in ≥10% of cortical PTCs (short arrows) with 3 to 4 in the most severely involved PTC (long arrow). Please note the slightly distended, open appearance of the capillary which can be a helpful feature; PAS, original magnification ×400. B, Banff Lesion Score ptc2—Moderate peritubular capillaritis defined as at least 1 cell in ≥10% of cortical PTCs (short arrows) with 5-10 in most severely involved PTC (long arrow); PAS, original magnification ×400. C, Banff Lesion Score ptc3—Severe peritubular capillaritis defined as at least 1 cell in ≥10% of cortical PTCs (short arrows) with >10 in most severely involved PTC (long arrow). PAS, original magnification ×400. D, This peritubular capillary is cut longitudinally (short arrow) and although containing 4 mononuclear cells is to be disregarded for scoring. However, the neighboring peritubular capillary (long arrow) is cut orthogonally and would qualify for Banff Lesion Score ptc1 provided that at least 10% of all PTCs contain at least 1 leukocyte. PAS, original magnification ×400.
FIGURE 8. Banff Lesion Score C4d. A, IHC staining with peroxidase yielding a brown reaction product for C4d. An example of C4d3, this image demonstrates linear and circumferential staining of endothelial cells in virtually all peritubular capillaries. The staining was similar in all areas of the cortex and the medulla. The proportion of stained peritubular capillaries and medullary vasa recta informs the score. B, IF staining for C4d. This image shows an example of a Banff Lesion Score of C4d3; using IF, a minimum score of C4d ≥ 2 is considered positive. In addition to this, the staining intensity for an individual capillary or medullary vas recta must be at least 1+ on the usual scale from negative, trace, 1+, 2+ to 3+. Indirect IF, mouse antihuman C4d followed by fluorescein isothiocyanate-conjugated anti-mouse IgG, original magnification ×100.

FIGURE 9. Banff Lesion Scores for ct (tubular atrophy) and ci (interstitial fibrosis). The ci and ct Scores are both based on calculating the total percentage of cortex involved and require a diligent assessment of all foci of ct and ci as this process is often multifocal; ct and ci scores may not always be equally advanced. A, This image demonstrates an area of nonatrophic tubules (long arrow), compared to an area of tubular atrophy (short arrow) without an obvious increase in interstitial fibrosis. PAS, original magnification ×200. There are different morphological types of tubular atrophy with differing histological appearances, including conventional, thyroidization, and endocrine-like types. B, Tubular atrophy of conventional type with interstitial fibrosis. Tubular areas are separated by areas of interstitial fibrosis and tubules show thickened basement membranes and > 50% reduction in tubular diameter (long arrows). PAS, original magnification ×200. C, Thyroidization type atrophy. Here, tubules appear dilated, have flattened epithelial cells, and contain eosinophilic and brightly periodic-acid-Schiff-positive uromodulin casts (long arrow). PAS, original magnification ×200. D, Endocrine-like type, characterized by shrunken tubules with cuboidal epithelium and “tubular simplification” (long arrow). Compared with the other types of tubular atrophy, endocrine-like type does not have thickened basement membranes but still counts toward the ct score. PAS, original magnification ×400.
Chronic Active AMR, but can also be seen in association with thrombotic microangiopathy of other causes than AMR, e.g., hepatitis C virus infection, hypertensive glomerulopathy, and glomerulonephritis. In analogy to Banff Lesion Score g, even in the presence of an explanation other than rejection for GBM double contours, Banff Lesion Score cg shall still be applied. Banff Lesion Score cg is not scored in ischemic or segmentally sclerosed glomeruli. Late ischemic glomerulopathy is defined as “thickening, wrinkling and collapse of glomerular capillary walls associated with

FIGURE 10. Visual analog scales provided by the Banff Working Group on Fibrosis. This working group developed schematic diagrams to facilitate and standardize scoring of Banff Lesion Scores ci and ct. A, Scale for the assessment of interstitial fibrosis without tubular atrophy. B, Scale for the assessment of diffuse tubular atrophy with “replacement fibrosis.” Reproduced with kind permission from American Journal of Transplantation.

FIGURE 11. More visual analog scales provided by the Banff Working Group on Fibrosis. A, Scale for the assessment of patchy (left) and confluent (right) interstitial fibrosis without glomeruli. B, Scale for patchy (left) and confluent (right) fibrosis with glomeruli. Reproduced with kind permission from American Journal of Transplantation.
extracapillary fibrotic material. As stated above, the earliest lesion of TG (cg1a) requires transmission EM for diagnosis. To detect such lesions, it is recommended that at centers with EM capability, “ultrastructural studies should be performed in all biopsies from patients who are sensitized, have documented DSA at any time posttransplantation and/or who have had a prior biopsy showing C4d staining, glomerulitis and/or peritubular capillaritis”. It is also advised that EM be considered in all biopsies performed from 6 months posttransplantation onward and in for-cause biopsies done from 3 months posttransplantation onward to determine if early changes of TG are present, prompting testing for DSA. Electron microscopy is also recommended for any biopsy done for the indication of increasing or new onset proteinuria.

cg0—No GBM double contours by light microscopy (LM) or EM.

cg1a—No GBM double contours by LM but GBM double contours (incomplete or circumferential) in at least 3 glomerular capillaries by EM, with associated endothelial swelling and/or subendothelial electron-lucent widening.

cg1b—Double contours of the GBM in 1-25% of capillary loops in the most affected nonsclerotic glomerulus by LM; EM confirmation is recommended if EM is available.

cg2—Double contours affecting 26 to 50% of peripheral capillary loops in the most affected—glomerulus.

cg3—Double contours affecting more than 50% of peripheral capillary loops in the most affected-glomerulus.

Banff Lesion Score mm (Mesangial Matrix Expansion)

This score evaluates the percentage of glomeruli with “moderate mesangial matrix expansion” in relation to all nonsclerosed glomeruli. Banff 1997 defines moderate mesangial matrix increase as “expansion of the matrix in the mesangial interspace to exceed the width of 2 mesangial cells in the average in at least 2 glomerular lobules”. An example is shown in Figure 15. Banff Lesion Score mm is currently not used to reach a Diagnostic Category and is purely descriptive.

mm0—No more than mild mesangial matrix increase in any glomerulus.

mm1—At least moderate mesangial matrix increase in up to 25% of nonsclerotic glomeruli.

mm2—At least moderate mesangial matrix increase in 26% to 50% of nonsclerotic glomeruli.

mm3—At least moderate mesangial matrix increase in >50% of nonsclerotic glomeruli.

Banff Lesion Score ah (Arteriolar Hyalinosis)

This score evaluates the extent of arteriolar hyalinosis (Figure 16). The first edition of the Banff Classification defined ah as “nodular hyaline afferent arteriolar thickening
suggestive of cyclosporine toxicity”; however, in Banff 1997 and later updates, Banff Lesion Score $ab$ is defined simply as PAS-positive arteriolar hyaline thickening, as a finding of “uncertain significance”. An asterisk “*$*$” is added to the $ab$ score when arteriolitis is present (eg, $ab0*$, $ab2*$).5 Banff Lesion Score $ab$ is currently not used to reach a diagnostic category and is purely descriptive.

$ab0$—No PAS (PAS)-positive hyaline arteriolar thickening.

$ab1$—Mild to moderate PAS-positive hyaline thickening in at least 1 arteriole.

$ab2$—Moderate to severe PAS-positive hyaline thickening in more than 1 arteriole.

$ab3$—Severe PAS-positive hyaline thickening in many arterioles.11

**Banff Lesion Score $aah$ (Hyaline Arteriolar Thickening)**

This Banff Lesion Score provides an alternative way of quantifying arteriolar hyalnosis. It was proposed in the 2007 update, because of the insufficient reproducibility of the Banff Lesion Score $ab$.8 This alternative tries to reach better reproducibility by focusing on circumferential or noncircumferential hyalnosis and the number of arterioles. Still, this lesion cannot be considered specific, that is, diagnostic for calcineurin inhibitor-related arteriolopathy. The use of this Banff Lesion Score $aah$ has been left as optional since its introduction in 2007, no final decision has been reached whether it shall replace Banff Lesion Score $ab$. Banff Lesion Score $aah$ is currently not used to reach a diagnostic category and is purely descriptive.

$aah0$—No typical lesions of calcineurin inhibitor-related arteriolopathy.

$aah1$—Replacement of degenerated smooth muscle cells by hyaline deposits in only 1 arteriole, without circumferential involvement.

$aah2$—Replacement of degenerated smooth muscle cells by hyaline deposits in more than 1 arteriole, without circumferential involvement.

$aah3$—Replacement of degenerated smooth muscle cells by hyaline deposits with circumferential involvement, independent of the number of arterioles involved.11

**Banff Lesion Score $ti$ (Total Inflammation)**

This lesion score evaluates the extent of total cortical inflammation. According to the Banff 2007 update and in contrast to the Banff Lesion Score $i$, all of the cortical parenchyma, including areas of interstitial fibrosis and tubular atrophy (IFTA), subcapsular cortex and perivascular cortex including nodular infiltrates are considered for $ti$ scoring.8 Mengel et al found Banff Lesion Score $ti$ to be better predictive of poor graft outcomes than the Banff Lesion Score $i$ in cases where at least mild IFTA was present.20 The association between interstitial inflammation in areas of IFTA as reflected in Banff Lesion Score $i$-IFTA and decreased graft survival was noted by Mannon et al21 and subsequently confirmed by others.22,23 As a consequence, Banff Lesion Score $ti$ became part of the criteria for a diagnosis of Chronic Active TCMR Grade IA and IB;12 both Banff Lesion Scores $ti$ and i-IFTA must be at least 2 to consider a diagnosis of Chronic Active TCMR Grade IA or IB.12

$ti0$—No or trivial interstitial inflammation (<10% of total cortical parenchyma).

$ti1$—10-25% of total cortical parenchyma inflamed.

$ti2$—26-50% of total cortical parenchyma inflamed.

$ti3$—>50% of total cortical parenchyma inflamed.11

**Banff Lesion Score $i$-IFTA (Inflammation in Area of IFTA)**

This score evaluates the extent of inflammation in scarred cortex, ie, areas that qualify for Banff Lesion Scores $ci$ and $ct$ (Figure 17). The Banff Lesion Score $i$-IFTA was first introduced to the Banff Classification in 2015.11 Both Banff Lesion Scores $ti$ and $i$-IFTA must be at least 2 to consider a diagnosis of Chronic Active TCMR Grade IA or IB.12

$i$-IFTA0—No inflammation or less than 10% of scarred cortical parenchyma.

$i$-IFTA1—Inflammation in 10% to 25% of scarred cortical parenchyma.

$i$-IFTA2—Inflammation in 26% to 50% of scarred cortical parenchyma.

$i$-IFTA3—Inflammation in >50% of scarred cortical parenchyma.11

**BANFF DIAGNOSTIC CATEGORIES**

Table 1 presents the Banff Diagnostic Categories and is based on the original table of the most recent Banff update from 2017.12 Readers should stay alert to future updates on the Banff Foundation website (www.banfffoundation.org) informed by updates to the Banff Classification from 2019 onward.

**CRITICAL APPRAISAL**

Since 1991, the Banff classification has undergone several amendments, reflecting the growing body of knowledge in transplant pathology. These amendments have been based on a consensus reached at the biannual Banff meetings. This constant refinement based on emerging data is a strength of the Banff process and has led to the worldwide dominance
of the Banff Classification for diagnostic practice, research and clinical trials. However, the iterative fashion in which the definitions and rules were published has dispersed the relevant content and created ambiguities. This has led to the creation of the Banff Rules and Dissemination Working Group in the aftermath of the Banff Meeting in Barcelona in March 2017. The aim of the Working group is not to alter the content of the Banff Classification. Rather, it shall collate all relevant Banff content in a central repository under the auspices of the Banff Foundation for Allograft Pathology, with a single updatable content, similar to the Union for International Cancer Control’s TNM Classification. Changes in the content of the Banff Classification must only be made through review of evidence and expert consensus at the Banff meetings or within the relevant other Working Groups. Like the collation of content above, the following critical appraisal is based on this mission and does not touch on the content of the Banff Classification itself.

Although the Banff Lesion Scores required for a diagnosis of AMR have recently undergone a partial overhaul and a dedicated Working Group is reexamining the Banff Lesion Scores for TCMR, no or little effort has been devoted to the Additional Diagnostic Parameters in Table 3. For example, “Acute Tubular Injury In The Absence Of Any Other Cause” as a criterion for active AMR is as important as Banff Lesion Scores $v$, $g$ or $ptc$, yet this feature is still imperfectly defined, the last definition dating back to the 1995 update. Another example is “infection,” which precludes the use of Banff Lesion Score $ptc$ alone as a criterion for AMR. Use of the isolated term “infection” is ambiguous in the context of whether inflammation in the transplant should be considered as evidence for rejection or not. We would recommend treating these Additional Diagnostic Parameters like the Banff Lesion Scores, presenting them in clear and consistent wording, and, whenever necessary, by providing guidance through meaningful definitions elaborated over time through Working Groups and in alignment with the respective diagnostic criteria applied.

Among the Banff Lesion Scores, the Banff Lesion Score $cv$ has a confusing array of terminologies, appearances and diagnostic implications. “Arterial fibrointimal thickening” or “vascular fibrous intimal thickening” imply a chronic fibrous change, whereas arterial intimal thickening can be cellular and nonfibrous in “transplant vasculopathy” or “chronic
allograft arteriopathy”.

As a manifestation of chronic TCMR, it is defined as “arterial intimal fibrosis with mononuclear cell infiltration in fibrosis, formation of neointima” whereas, as a criterion for AMR chronicity, it is defined as “arterial intimal fibrosis of new onset, excluding other causes; leukocytes within the sclerotic intima favor chronic AMR if there is no prior history of biopsy-proven TCMR with arterial involvement but are not required.” In clinical practice, it might not always be possible to exclude prior TCMR or to precisely diagnose “Arterial intimal fibrosis of new onset” as a criterion for AMR chronicity. A related problem is attached to Banff Lesion Score cg; “evidence of chronic thrombotic microangiopathy (TMA)” excludes the use of Banff Lesion Score cg > 0 as a criterion for AMR chronicity, whereas Active AMR can be diagnosed with TMA, as long as it is “in the absence of any other cause [than AMR]”. Because Active AMR causing TMA can lead to glomerular lesion qualifying as TG, it would make sense to change the cg criterion to only exclude chronic TMA of any other cause than AMR.

The use of asterisks (“*”) attached to Banff Lesion Scores v, i, ah and ptc is problematic and widely neglected. Their reproducibility and diagnostic value are unknown, and they are ambiguous: an asterisk behind the Banff Lesion Score ptc signifies only mononuclear cells and absence of neutrophils, whereas the asterisk behind Banff Lesion Score i denotes a significant neutrophilic, eosinophilic or plasmacellular component in the infiltrate, and these different cell types can have widely differing implications. We suggest the Banff community should reassess these modifiers, either by improving their definitions and assigning them a significance or by abandoning them.

Inevitably, the Banff Classification has focused mainly on features of rejection, but with Banff Lesion Scores developed for other features with little or no guidance on their contribution to diagnosis. An example for this is Banff Lesion Score aah, originally intended to replace the poorly reproducible Banff lesion score ah. However, its use is still optional, and it has neither been widely adopted nor used in any of the Banff Diagnostic Categories. The Banff community should reassess arteriolar hyalinosis lesion scores, and clarify grading and diagnostic implications.

Regarding the Banff Diagnostic Categories, a clear diagnostic pathway should be recommended when dealing with Borderline or Acute TCMR (Banff Diagnostic Categories 3 and 4) in the presence of BK Virus Nephropathy, Pyelonephritis or other infectious diseases of the transplant, as well as AMR with glomerulitis in the presence of recurrent or de novo glomerulonephritis. These issues could be referred to the Banff TCMR and Glomerulonephritis Working Group respectively. The definition of Banff Borderline with regards to the Banff Lesion Score i threshold (i0 or i1) is still ambiguous but should be resolved by the TCMR Working Group.

There are uncertainties around the application of transmission EM in the diagnosis of AMR which are currently being addressed by the Electron Microscopy Working Group. These issues include precise guidelines for indications and methods for application of EM in transplant biopsies; perhaps also the introduction of a new Banff Lesion Score for multilamination of the basement membranes of peritubular capillaries which we have covered as an Additional Diagnostic Parameter for now.

Another critical issue is related to the molecular diagnostics of AMR and TCMR. Although the current Banff classification endorses the use of molecular diagnostics in the definition of AMR, there is limited guidance regarding methods and diagnostic cut-offs, which could be elaborated by the Molecular Working Group.

Lastly, the introduction of the new diagnostic categories of Chronic Active TCMR is likely to undergo changes informed by the TCMR Working Group. Before Banff 2017, there were no specific criteria for Chronic Active TCMR outside of arteries, and tubulitis was only scored in nonatrophic and mildly atrophic tubules, effectively excluding moderately and severely atrophic tubules. To avoid having 2 separate criteria for Banff Lesion Score t in Acute versus Chronic Active TCMR, it was decided that for both diagnoses tubulitis would be scored in all tubules except severely atrophic tubules. The difference between Banff 2017 and previous versions of the classification with respect to Acute TCMR is that tubulitis in moderately atrophic tubules is now counted toward Banff Lesion Score t. Because the latter was done for clarity and to avoid confusion rather than on the basis of specific evidence, it would be beneficial that future studies be done to address the most clinically relevant threshold for the level of atrophy permitted in scoreable tubules, especially for diagnosis of Acute TCMR. In addition, the 2017 changes to the TCMR criteria also suggest future work be aimed at examining the response of Chronic Active TCMR to steroids and other anti–T cell therapies (eg, thymoglobulin), determining if there are differences in this response between: (1) Grade IA versus Grade IB Chronic Active TCMR and (2) biopsies with Chronic Active TCMR that would otherwise meet
criteria for acute TCMR (ie, with Banff Lesion Score $i \geq 2$) and those that would not (with Banff Lesion Score $i \leq 1$). The alignment of diagnoses from the spectrum of Acute TCMR with those from the spectrum of Chronic Active TCMR of different compartments could be problematic. For example, a biopsy with Banff Lesion Score $v1$ fulfilling also the criteria for chronic active TCMR grade IB would be diagnosed as the latter only, as according to Banff 2017, a diagnosis of Chronic Active TCMR precludes the diagnosis even of higher grade Acute TCMR. In such cases, however, the use of modifying text independent from Banff diagnostic categories should be considered (eg, Acute TCMR

![Figure 16](image)

**FIGURE 16.** Banff Lesion Score $ah$ (arteriolar hyalinosis). A, Banff Lesion Score $ah1$—mild focal arteriolar hyalinosis (arrow). PAS, original magnification $\times 630$. B, $ah2$—Moderate arteriolar hyalinosis (arrow). PAS, original magnification $\times 630$. C, Banff Lesion Score $ah2$—Note in this image there is both linear (short arrow) and nodular hyalinosis (long arrow). For a score of $ah2$, more than 1 arteriole displaying moderate to severe is required. Jones silver stain, original magnification $\times 630$. D, Banff Lesion Score $ah3$—severe circumferential arteriolar hyalinosis with luminal occlusion. For Banff Lesion Score $ah3$, hyalinosis of this severity (arrow) must be present in many arterioles as depicted here. PAS, original magnification $\times 630$.

![Figure 17](image)

**FIGURE 17.** Banff Lesion Score $i$-IFTA (Inflammation in areas of IFTA). Image A shows Inflammation in areas of IFTA (arrow). This Lesion Score ranges from 0 to 3, based on the percentage of scarred areas of the cortex (ie, areas qualifying for $ci$ and $ct$) involved by inflammation. It is one of the criteria necessary for a diagnosis of Chronic Active TCMR Grade IA or IB. Masson trichrome, original magnification $\times 200$. B, In contrast shows interstitial fibrosis without significant infiltrate (arrow). H&E, original magnification $\times 400$. 

© 2018 Wolters Kluwer Roufosse et al
Grade II with a chronic active tubulointerstitial component; Acute TCMR Grade II with isolated intimal arteritis [isolated v].

**PROSPECTS**

Although this article is intended to provide a comprehensive and convenient desk-top reference, it is destined to expire with the publication of the 2019 Banff update. After this update, a Web resource will serve as the continuously updated go-to resource for the relevant Banff content. Depending on the progress in the definitions and diagnostic rule sets we are aiming to develop web-based resources such as diagnostic algorithms to further strengthen standardization and reproducibility of the Banff Classification for clinical practice and

### TABLE 3.

These additional diagnostic parameters, some histopathologic, some clinical, are derived from the diagnostic algorithms in Table 1

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Required for diagnostic category</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute Thrombotic Microangiopathy In The Absence Of Any Other Cause (Figure 18)</td>
<td>Active and Chronic Active AMR</td>
</tr>
<tr>
<td>Acute Tubular Injury In The Absence Of Any Other Apparent Cause</td>
<td>Active AMR</td>
</tr>
<tr>
<td>Absence Of Recurrent Or De Novo Glomerulonephritis</td>
<td>Active and Chronic Active AMR</td>
</tr>
<tr>
<td>Infection</td>
<td>Active and Chronic Active AMR</td>
</tr>
<tr>
<td>Arterial Intimal Fibrosis Of New Onset, Excluding Other Causes</td>
<td>Chronic AMR and Chronic Active AMR</td>
</tr>
<tr>
<td>Leukocytes Within The Sclerotic (Arterial) Intima Favor Chronic AMR</td>
<td>Chronic Active AMR and Chronic AMR</td>
</tr>
<tr>
<td>Increased Expression Of Thoroughly Validated Gene Transcripts/Classifiers In The Biopsy Tissue Strongly Associated With AMR</td>
<td>Active AMR, Chronic Active AMR, Chronic AMR</td>
</tr>
<tr>
<td>Severe Peritubular Capillary Basement Membrane Multilayering (Figure 19)</td>
<td>Chronic AMR and Chronic Active AMR, Chronic Active TCMR Grade IA and IB</td>
</tr>
<tr>
<td>Arterial Intimal Fibrosis With Mononuclear Cell Inflammation In Fibrosis And Formation Of Neointima</td>
<td>Chronic Active TCMR Grade II</td>
</tr>
<tr>
<td>Prior Evidence Of DSA</td>
<td>Chronic AMR</td>
</tr>
<tr>
<td>Serologic Evidence Of DSAs (DSA To HLA Or Other Antigens)</td>
<td>Active AMR, Chronic Active AMR, Chronic AMR</td>
</tr>
<tr>
<td>Prior Documented Diagnosis Of Active Or Chronic Active AMR</td>
<td>Chronic AMR</td>
</tr>
<tr>
<td>Prior History Of TCMR</td>
<td>Chronic Active AMR and Chronic AMR</td>
</tr>
<tr>
<td>Evidence Of Chronic TMA</td>
<td>Chronic Active AMR and Chronic AMR</td>
</tr>
<tr>
<td>C4d Staining On Fresh-Frozen Or Paraffin-Embedded Tissue</td>
<td>C4d Staining Without Evidence Of Rejection, Active AMR, Chronic AMR</td>
</tr>
<tr>
<td>BK-Virus Nephropathy</td>
<td>Other Changes Not Considered To Be Caused By Acute Or Chronic Rejection</td>
</tr>
<tr>
<td>Posttransplant Lymphoproliferative Disorder</td>
<td></td>
</tr>
<tr>
<td>Calcineurin Inhibitor Toxicity</td>
<td></td>
</tr>
<tr>
<td>Acute Tubular Injury</td>
<td></td>
</tr>
<tr>
<td>Recurrent Disease</td>
<td></td>
</tr>
<tr>
<td>De Novo Glomerulopathy (Other Than TG)</td>
<td></td>
</tr>
<tr>
<td>Pyelonephritis</td>
<td></td>
</tr>
<tr>
<td>Drug-Induced Interstitial Nephritis</td>
<td></td>
</tr>
</tbody>
</table>

Depending on the constellation of findings they may be required in addition to the Banff Lesion Scores to determine the Banff Diagnostic Categories.

**FIGURE 18.** Acute TMA. A, An acute TMA affecting a glomerulus with fibrin thrombi (long arrows) and fragmented red blood cells (short arrow) in capillary loops. Trichrome, original magnification ×400. B, An acute TMA affecting a small arteriole (arrow). Acute TMA is one of the histological features used as histological evidence of acute tissue injury in Active AMR. However, TMA is not specific for AMR and can be seen in, for example, recurrent disease or calcineurin inhibitor toxicity. Trichrome, original magnification ×400.
research. It should be emphasized that the Banff Classification of Kidney Allograft Pathology does not cover all relevant aspects of transplantation medicine. Allograft transplantation only reaches 10% of patients needing new organs. Through regenerative medicine and tissue engineering and other optimizing initiatives we will eventually be able to provide organs to everyone in need. For this, we will need a new Banff Classification of Tissue Engineering Pathology.24,25

FIGURE 19. Severe Peritubular Capillary Basement Membrane Multilayering (PTCML) as demonstrated by EM. A, This Additional Diagnostic Parameter is a criterion for AMR chronicity. It is defined as 7 or more layers of basement membrane in at least 2 additional capillaries. This particular capillary shows 8 layers (arrow). Transmission EM, original magnification ×14000. B, This image demonstrates a peritubular capillary with 5 layers of basement membrane (arrow). Transmission EM, original magnification ×10000.

FIGURE 20. Banff Classification Diagnostic Category 6 (other). These images illustrate some of the more common examples of key lesions specified under category 6. A, Pyelonephritis with neutrophilic casts (arrow) and neutrophilic infiltrates with tubulitis. H&E, original magnification ×200. B, BK virus nephropathy with typical ground glass intranuclear inclusions as seen on hematoxylin and eosin stain (arrows). H&E, original magnification ×400. C, Acute tubular injury with widespread isometric vacuolization of tubular epithelial cells (arrow) associated with acute Calcineurin Inhibitor Toxicity and other forms of injury. H&E, original magnification ×400. D, Recurrent glomerulonephritis (membranoproliferative immune complex glomerulonephritis type I in this case) with split GBMs (arrow). The diagnosis was confirmed and TG excluded by positive IF for immunoglobulin heavy-, light-chains and complement slit products as well as abundant subendothelial electron dense immune complex deposits on EM. PAS, original magnification ×400.
reflecting the new challenges of delivering the right cells to the right places in a bioengineered organ and having them function normally. Rejection will no longer be the primary threat in bioengineered organs. For a decade or more the new Banff Classification of Tissue Engineering Pathology will be used concurrently with the existing Banff Classification of Allograft Pathology.

Getting the right cells in the right places sounds simple, but in fact, we have poor knowledge of what all the normal cell types in transplanted organs are. For instance, in the kidney, we have traditionally taught that there are 26 cell types, but in fact, high throughput single cell analysis in the Human Cell Atlas Project shows many more than that and can determine not only cell identity but also lineage and activation state. The transplantation and transplantation pathology community need to embrace Human Cell Atlas technology, so we are not blindsided by this new technology. The scale of the likely impact of the Human Cell Atlas Project on nephrology and transplantation is currently being analyzed (Moghe I, Magor B, and Solez K, article in preparation, 2018).

ACKNOWLEDGMENTS

The authors would like to acknowledge the help in the preparation of the visual analog scales from Christopher Bellamy, Alton “Brad” Farris and Daniel Serón.

REFERENCES