Delayed Cerebral Ischemia in Aneurysmal Subarachnoid Hemorrhage: Proposal of an Evidence-Based Combined Clinical and Imaging Reference Standard

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ABSTRACT

SUMMARY: Aneurysmal subarachnoid hemorrhage is associated with high morbidity and mortality, with delayed neurologic deficits from delayed cerebral ischemia contributing to a large portion of the adverse outcomes in this patient population. There is currently no consensus reference standard for establishing the diagnosis of delayed cerebral ischemia either in the research or clinical settings, ultimately limiting strategies for preventing delayed infarction and permanent neurologic deficits. There are currently both clinical and imaging-based criteria for the diagnosis of delayed neurologic deficits and vasospasm, respectively, however, neither clinical nor angiographic assessment alone has been shown to identify patients who develop adverse outcomes from delayed infarction. Thus, the purpose of this work is to propose a 3-tiered combined imaging and clinical reference standard based on evidence from the literature to standardize the diagnosis of delayed cerebral ischemia, both to allow consistency across research studies and to ultimately improve outcomes in the clinical setting.

ABBREVIATIONS: aSAH = aneurysmal subarachnoid hemorrhage; DCI = delayed cerebral ischemia; MRP = MR perfusion

Aneurysmal subarachnoid hemorrhage (aSAH) is associated with high morbidity and mortality.1,2 The first 2 weeks following aSAH are critical in the management of these patients because they are prone to develop several life-threatening complications, including delayed neurologic deficits,3 which often arise from delayed cerebral ischemia (DCI), a major contributor to the adverse outcomes in this population.3-5 Delayed cerebral ischemia manifests in approximately 30% of patients with aSAH and typically occurs between days 4 and 9 after the initial hemorrhage, though it can range from 3 to 14 days.

There remains a lack of standard criteria for defining DCI in the clinical setting,3,6,7 with a recent literature review describing at least 8 terms to define the concept of DCI in aSAH.6 Debate over the role of clinical and imaging assessments in defining DCI has occurred for both clinical and research purposes.3,6,8-10 For example, although the terms “DCI” and “vasospasm” have been used interchangeably, attempts have been made to distinguish DCI from vasospasm, with the former often determined clinically, and the latter, radiographically,6 because not all patients with clinical neurologic deficits have angiographic vasospasm and not all patients with angiographic vasospasm have neurologic deficits that correspond to the arterial territory of vasospasm.11,12 Additionally, while severe vasospasm may cause decreased cerebral perfusion, a substantial percentage of patients develop infarction without evidence of vasospasm, suggesting that DCI should be defined as a pathologic process, of which vasospasm may represent a contributing factor.12,13

Thus, the aim of this article is to propose an evidence-based reference standard for DCI that incorporates both clinical assessments of neurologic deterioration and imaging assessments of vasospasm, perfusion deficits, and infarction to provide a consistent, uniform standard across a wide range of clinical and research applications. The classification of levels of evidence supporting this reference standard is based on the Levels of Evidence criteria proposed by the Oxford Centre of Evidence Based Medicine (www.cebm.net).14 Two independent reviewers assessed levels of evidence for each tier, and in the case of discordance, evidence level assignments were made by consensus.
DESCRIPTION OF THE COMBINED CLINICAL AND IMAGING REFERENCE STANDARD

Primary Level: Outcome-Based Criteria

Summary. The primary level classifies patients as having DCI if a new infarction on imaging or new permanent neurologic deficit develops. A new infarction on imaging is determined on CT or MR imaging within 6 weeks after aSAH ictus that was not present on imaging up to 48 hours after aneurysm occlusion and was not attributable to other causes such as surgical clipping, endovascular treatment, ventricular catheter placement, intraparenchymal hematoma, or cerebral herniation. A new permanent neurologic deficit is determined on clinical examination as a new neurologic deficit distinct from the baseline examination performed immediately after aneurysm rupture or aneurysm occlusion and not attributable to other causes. Baseline neurologic examination must be considered after full cardiorespiratory, hemodynamic, and metabolic resuscitation as well as treatment of other factors such as seizures and hydrocephalus. Patients who do not meet either criterion are referred to the secondary level, as described in a subsequent section.

Evidence: Level 1A evidence exists to support these proposed outcomes-based criteria for determining DCI.

An ideal reference standard should reliably identify patients with a high risk of poor outcomes who may benefit from intervention. In large prospective cohort studies, the greatest predictors of severe disability or death at 3 months were a new focal neurologic deficit, a new infarction on follow-up imaging, or both.6,15,16 Additionally, a large systematic review and meta-analysis of all randomized placebo-controlled trials evaluating the efficacy of protective strategies in aSAH concluded that a reduced incidence of cerebral infarction is significantly associated with improved functional outcome.17 In fact, new cerebral infarction alone was as strongly correlated with poor 3-month functional outcome as the combination of a new neurologic deficit and corresponding ischemic changes on follow-up neuroimaging.6 Furthermore, cerebral infarction on noncontrast CT was the primary outcome measure in the early trials of nimodipine, an agent with strong evidence for neuroprotection of DCI.18

FIG 1. Proposed multitiered reference standard in DCI. Three-tiered DCI reference standard algorithm, ordered from top to bottom. Asterisk indicates neuromonitoring devices such as cerebral microdialysis and oximetry. Double asterisks indicate whether the reference standard is used for clinical assessment and treatment decisions based on the risk/benefit ratio for treatment. If there is low risk, treatment for DCI is recommended. If there is high risk, the patient should re-enter the algorithm.

Reference standard for DCI

A-SAH patients

Primary level

DCI-related Outcomes

OUTCOMES CRITERIA: Clinical and/or Imaging

YES

DCI

NO

Secondary level

DCI related to Vasospasm

CORRELATION OF CLINICAL AND IMAGING CRITERIA: TCD, CTA, MRA, DSA

YES

DCI

Clinical and Imaging data both Positive

NO

Clinical and Imaging data both Negative

Tertiary level

Asymptomatic or Poor clinical exam

CORRELATION WITH PERFUSION STUDY (e.g. CTP, MRP, Neuromonitoring*)

POSITIVE

DCI

NEGATIVE

INSUFFICIENT EVIDENCE FOR DCI**

CORRELATION WITH PERFUSION STUDY (e.g. CTP, MRP, Neuromonitoring*)

POSITIVE

DCI

NEGATIVE

Both clinical and initial imaging exams support a diagnosis of DCI related to vasospasm.

Both clinical and initial imaging exams do NOT support a diagnosis of DCI related to vasospasm.

Permanent neurologic deficit or cerebral infarction on follow-up CT or MRI.
While angiographic vasospasm has traditionally been the primary focus of interventions and prediction of outcomes, the lack of evidence demonstrating improved outcomes with vasospasm prevention\(^3,13\) has led to incorporating this criterion combined with clinical correlation in the secondary level below.

**Secondary Level: Correlation of Clinical and Vascular Imaging Criteria**

**Summary.** The secondary level classifies patients as having DCI if both clinical deterioration and angiographic vasospasm occur. Clinical deterioration is determined by bedside examination and comprises the development of a new neurologic deficit (such as hemiparesis, hemiplegia, aphasia, depressed consciousness, and so forth), a decrease of at least 2 points on the Glasgow Coma Scale, or a decrease of at least 1 point in the motor score, lasting >1 hour at any point after aneurysm occlusion and not attributable to other causes. Vascular imaging for the evaluation of vasospasm includes imaging modalities, such as transcranial Doppler sonography, CTA, MRA, and DSA. Patients with neurologic deterioration and 1 imaging test supporting a diagnosis of vasospasm are classified as having DCI. On the other hand, patients without neurologic deterioration and 1 imaging test without findings of vasospasm are classified as not having DCI. However, patients with either positive clinical or imaging findings that do not correlate with each other are referred to the tertiary level, as described in a subsequent section.

Evidence: Level 1B evidence exists to support using clinical and vascular imaging data for determining DCI.

Evaluation of patients for DCI at the secondary level is most valuable in the clinical setting at the point of care when treatment decisions are made. The primary goal of treatment is to prevent cerebral infarction and permanent neurologic deficits. Thus, traditionally, imaging assessment of vasospasm has been used as a surrogate marker to assist in the diagnosis of DCI, especially given that neurologic deterioration is poorly evaluated in sedated or obtunded patients. Angiographic vasospasm, seen on DSA or CTA, is perhaps the most commonly used surrogate imaging marker in this patient population. Vasospasm has been shown to be strongly associated with DCI, cerebral infarction, poor outcome, and increased mortality within several retrospective and prospective cohort studies, including a post hoc analysis of data from the CONSCIOUS (Clazosentan to Overcome Neurological Ischemia and Infarct Occurring after Subarachnoid Hemorrhage)-1 trial.\(^{11,20-23}\) However, an analysis of data from 2 systematic reviews and a post hoc analysis did not demonstrate an improvement in outcome with a reduction in angiographic vasospasm.\(^9\) Evidence from both prospective and retrospective cohort studies suggests that patients with angiographic vasospasm and correlated symptoms have worse hospital complications and subsequent disability compared with angiographic vasospasm alone.\(^5,16\) However, there is less evidence to demonstrate the prognostic importance of angiographic vasospasm correlated with symptoms, thus placing this criterion at the secondary level. Although relatively inferior in terms of sensitivity and specificity, transcranial Doppler sonography evaluations of the intracranial vessels can also be performed at bedside to identify arterial narrowing in patients who may be too unstable for more advanced angiographic techniques such as CTA, MRA, or DSA.\(^{24,25}\)

**Tertiary Level: Correlation of Physiologic Data with Clinical or Imaging Criteria**

**Summary.** The tertiary level classifies patients as having DCI if physiologic data correlates with either clinical deterioration or vasospasm. Patients with either clinical deterioration or vasospasm alone may undergo additional physiologic assessment of cerebral hemodynamics, either in the form of imaging such as CTP and MR perfusion (MRP) or neuromonitoring devices such as cerebral blood flow, oxygen tension monitoring, and cerebral microdialysis. Patients with findings suggestive of regional cerebral hypoperfusion or hypoxia that correlate with either clinical deterioration or vasospasm are classified as having DCI. Patients with clinical deterioration or vasospasm but normal physiologic data do not have sufficient evidence to be classified as having DCI.

Evidence: Levels of evidence to support using physiologic data for determining DCI range from 2A to 3B, depending on the technique.

While there is at least moderate evidence supporting the importance of symptomatic vasospasm in DCI at the secondary level, the importance of isolated image-based diagnoses of vasospasm in the absence of clinical findings is somewhat controversial, especially in the absence of infarction. However, a subset of patients with asymptomatic vasospasm will develop asymptomatic ischemia and subsequent infarction. A large prospective cohort identified asymptomatic infarction in approximately 20% of patients with aSAH, and furthermore, these patients had a higher frequency of death and moderate-to-severe disability at 3 months relative to patients with symptomatic infarction.\(^{26}\) Thus, there may be a subset of patients with apparently asymptomatic vasospasm who are at high risk of eventually developing clinical evidence of DCI, especially those who are comatose or have a ventriculostomy catheter, small-volume aSAH, or ischemia in noneloquent brain.\(^{26,27}\) —all representing complicating factors that are not infrequently encountered in the intensive care setting. Identifying this high-risk subset of patients with asymptomatic vasospasm may prompt measures to implement therapies to prevent the eventual development of DCI.

Conversely, the identification of patients with DCI and clinical deterioration in the absence of vasospasm poses a different important diagnostic challenge. While neurologic deterioration is likely multifactorial in these patients, a subset will go on to develop infarction without vasospasm. A retrospective study of infarction patterns in patients with aSAH found that approximately 17% of patients developed infarcts without imaging evidence of vasospasm, and even in patients with imaging positive for vasospasm, infarcts also developed in areas away from the vasospastic territories.\(^{28}\) Thus, this level in the algorithm would attempt to identify ischemia in patients with asymptomatic vasospasm or neurologic deterioration without evidence of large-vessel vasospasm.

Perfusion imaging such as CTP and MRP or less common modalities such as xenon-CT provide physiologic imaging assessments of cerebral hypoperfusion and ischemia that could identify patients at risk for infarction. In a retrospective cohort of 96 pa-
patients with aSAH, new CTP deficits seen as prolonged MTT and reduced CBF were significantly associated with subsequent infarction and permanent neurologic deficits. A smaller prospective study evaluating the test characteristics of CTP, CTA, and noncontrast CT obtained at baseline and after the onset of clinical deterioration determined that CTP had the best test performance for the subsequent diagnosis of DCI at discharge. Subsequently, systematic reviews evaluating CTP in aSAH within the broader context of diagnosing vasospasm and DCI found that relative CBF and MTT values correlated highly with subsequent DCI. Thus, there is level 2A evidence to support the role of CTP in the diagnosis of DCI.

Evidence to support the use of other imaging modalities to evaluate DCI is more limited. There are limited data evaluating the role of MRP in DCI; however, several small prospective cohort studies demonstrated that CTP, particularly CBF, correlates with MRP-derived values in the same patients within a close time interval, suggesting that MR imaging could also be used in this setting in case CTP is not performed. The data for the use of xenon-CT in DCI are even more limited; however, a small prospective cohort study in patients with poor-grade aSAH found that CBF reduction on xenon-CT was only moderately predictive of infarction in these patients and that not all reductions in CBF correlated with functionally relevant outcomes. Because these criteria can, in some cases, be evaluated before development of infarction and functional disability (ie, a stage in which impending DCI is still preventable), classification of patients with DCI at this level should theoretically provide maximal benefit from treatment.

A limitation of the secondary level is that patients without new neurologic deficits and without angiographic vasospasm can be misclassified as having no DCI. Comatose or heavily sedated patients have limited clinical assessment and may have suboptimal imaging, resulting in false-negatives for DCI. Although the agreement of clinical and imaging findings improves the specificity for identifying patients with DCI, given that neurologic assessment in patients with aSAH can be challenging and angiographic vasospasm does not necessarily correlate with DCI.

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**Tertiary Level.** The main strength of the tertiary level is improving the sensitivity of the DCI diagnosis by further evaluating discordant clinical and imaging findings from the secondary level, such as in patients with asymptomatic vasospasm or neurologic decline without angiographic vasospasm. Most important, this level allows further evaluation of comatose patients with suboptimal clinical assessments who have angiographic vasospasm as well as symptomatic patients who have suboptimal imaging. These patients often have worse outcomes in comparison with patients with symptomatic DCI, possibly related to delayed treatment. At this level, all patients undergo physiologic assessment of cerebral perfusion and hypoxia to correlate with either clinical or imaging findings suggesting DCI. Thus, this level will include patients who may have been excluded from the diagnosis due to lack of sufficient evidence at the other 2 levels.

A potential limitation of the tertiary level is that the breadth of modalities used to assess ischemia—ranging from noninvasive imaging to invasive tissue monitoring—has variable strength of...
evidence to support their use. From an imaging standpoint, CTP has the strongest evidence to support its use in diagnosing DCI; clinically, cerebral microdialysis has some evidence to support its use despite inconclusive results from a systematic review of the literature. There is limited evidence to support the use of the remaining modalities in diagnosing DCI in patients with aSAH.

Future Directions
While there is no perfect reference standard for this complex disease process, this multitiered algorithm attempts to capture the complexity of clinical and imaging findings in DCI according to evidence-based criteria. Specificity is emphasized in this multitiered reference standard with respect to evidence-based clinically relevant outcomes at the primary level, which are particularly valuable in the research setting to potentially improve translation of research findings into clinical practice. Most important, this reference standard approach also incorporates levels of evidence with greater sensitivity for use in clinical settings. The model is heavily weighted toward criteria with supportive statistical evidence and, through a multitiered algorithm, aims to limit the heterogeneity and controversy in defining DCI for research and, potentially, clinical application, combining both imaging and clinical assessments in the determination of DCI. The future direction for validation of this proposed reference standard through prospective studies may help to move forward both clinical care and research in this field.


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