



Published in final edited form as:

Am J Ophthalmol. 2021 August ; 228: 152–158. doi:10.1016/j.ajo.2021.03.043.

Classification criteria for multifocal choroiditis with panuveitis

The Standardization of Uveitis Nomenclature (SUN) Working Group^{*,1,2,3}

Abstract

Purpose: To determine classification criteria for multifocal choroiditis with panuveitis (MFCPU)

Design: Machine learning of cases with MFCPU and 8 other posterior uveitides.

Methods: Cases of posterior uveitides were collected in an informatics-designed preliminary database, and a final database was constructed of cases achieving supermajority agreement on diagnosis, using formal consensus techniques. Cases were split into a training set and a validation set. Machine learning using multinomial logistic regression was used on the training set to determine a parsimonious set of criteria that minimized the misclassification rate among the posterior uveitides. The resulting criteria were evaluated on the validation set.

Results: One thousand sixty-eight cases of posterior uveitides, including 138 cases of MFCPU, were evaluated by machine learning. Key criteria for MFCPU included: 1) multifocal choroiditis with the predominant lesions size >125 μm in diameter; 2) lesions outside the posterior pole

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³**Conflict of Interest:** Douglas A. Jabs: none; Antoine P. Brezin: none; Ralph D. Levinson: none; Peter McCluskey: none; Neal Oden: none; Alan G. Palestine: none; Russell W. Read: none; Jennifer E. Thorne: Dr. Thorne engaged in a portion of this research as a consultant and was compensated for the consulting service; Brett E. Trusko: none; Albert Vitale: none; Susan E. Wittenberg: none.

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(with or without posterior involvement); and either 3) punched-out atrophic chorioretinal scars or 4) more than minimal mild anterior chamber and/or vitreous inflammation. Overall accuracy for posterior uveitides was 93.9% in the training set and 98.0% (95% confidence interval 94.3, 99.3) in the validation set. The misclassification rates for MFPCU were 15% in the training set and 0% in the validation set.

Conclusions: The criteria for MFPCU had a reasonably low misclassification rate and appeared to perform sufficiently well for use in clinical and translational research.

PRECIS

Using a formalized approach to developing classification criteria, including informatics-based case collection, consensus-technique-based case selection, and machine learning, classification criteria for multifocal choroiditis with panuveitis were developed. Key criteria included multifocal choroiditis either with characteristic punched-out, atrophic scars and/or more than minimal vitreous inflammation. The resulting classification criteria had a low misclassification rate.

In 1984 Dreyer and Gass described a new posterior uveitic disease, multifocal choroiditis with panuveitis (MFPCU).¹ The disease had a retinal picture similar to that of the presumed ocular histoplasmosis syndrome in that there were “punched-out atrophic” chorioretinal scars of variable size, but differed in that there was a variable anterior chamber and vitreous inflammation and there was no evidence of prior histoplasmosis infection on serologic testing, skin testing, and chest radiography.¹ Most cases were bilateral. There often were lesions of variable character, including “active lesions” described as yellow to yellow-white, round or oval, sometimes irregular, and mildly elevated, with “punched-out atrophic” chorioretinal scars with variable hyperpigmentation at the edges. Lesions typically were >250µm in size.^{1,2}

Multifocal choroiditis with panuveitis is an uncommon uveitic disease. Most data on the disease come from case series.²⁻⁵ In one 6-year series of all patients with uveitis seen at a single, tertiary-care uveitis center in Australia, MFPCU accounted for 2.4% of all uveitic cases.⁶ The incidence of MFPCU has been estimated at 0.03 cases per 100,000 population per year.⁷ Multifocal choroiditis with panuveitis has been reported with a wide age range, but most cases occurred in young to middle-aged adults. Although MFPCU occurs in both men and women, the majority of reported cases have been in women. It has been reported in multiple ethnic groups, but the majority of cases appear to occur in Caucasians.²⁻⁵

The etiology of MFPCU is unknown, and it is unassociated with a systemic disease. The differential diagnosis of MFPCU includes those diseases which can produce a multifocal choroidopathy, such as punctate inner choroiditis (PIC), syphilis, tuberculosis (TB) in endemic areas, and sarcoidosis. Rarely, late-stage, untreated birdshot chorioretinitis (BSCR) may have a similar appearance. Occasionally serpiginous choroiditis not adjacent to the disc may look like MFPCU, but the characteristic fluorescein angiogram of serpiginous choroiditis typically allows the correct diagnosis to be made.^{2-5,8}

The active lesions of MFPCU have been described as yellow-orange round or oval, sometimes elevated, and typically >250 µm in size. The “punched-out” atrophic scars involve loss of choroid and retinal pigment epithelium in a circular fashion, typically with

pigment clumping at the edge. Fluorescein angiography of active MFCPU lesions has been reported as demonstrating multiple chorioretinal spots with progressive hyperfluorescence throughout the angiogram. By contrast, the atrophic scars demonstrate window defects on fluorescein angiography. Occasionally it can be difficult to differentiate choroidal neovascularization from an active MFCPU lesion of fluorescein angiography alone.⁸ On indocyanine green angiography, active MFCPU lesions have been reported to present as hypofluorescent spots that fade by the late phases of the angiogram, suggesting that the lesions are at the level of the choriocapillaris and/or retinal pigment epithelium.⁹ Fundus autofluorescence imaging has been reported as useful in assessing the activity of MFCPU lesions. Atrophic scars are hypo-autofluorescent, whereas active lesions are mildly hyper-autofluorescent.^{10–13} More lesions may be visible on fundus autofluorescent imaging than are evident clinically.¹³ Optical coherence tomography is useful in diagnosing macular edema and active choroidal neovascularization. It also has been reported to distinguish active choroidal lesions from atrophic scars. Optical coherence tomography angiography, although not routinely used, has been reported to differentiate choroidal neovascularization from active MFCPU lesions by detecting the abnormal subretinal vessels.¹⁴

Reported structural complications include macular edema, choroidal neovascularization, optic neuropathy, epiretinal membranes, and cataract.^{2,9,10,15,16} Choroidal neovascularization has been reported as the most common cause of vision loss.¹⁵ Incidences of visual impairment (20/50 or worse) and blindness (20/200 or worse) in involved eyes have been estimated at 0.19/eye-year (EY) and 0.12/EY, respectively, and in the better-seeing eye at 0.07/EY and 0.04/EY, respectively.¹⁵ High-dose oral corticosteroids have been reported to control the inflammation and decrease the occurrence of retinal structural complications, but doses low enough for long-term use (<10 mg/day) appear to be largely ineffective.^{15,17} Conversely, immunosuppression has been reported to reduce the occurrence of structural complications by over 80%.^{15,17} Hence, if treatment is needed, oral corticosteroids and immunosuppression appear to be the preferred approach.^{15,18} Choroidal neovascularization typically is treated with adjunctive anti-VEGF therapy.¹⁰

The Standardization of Uveitis Nomenclature (SUN) Working Group is an international collaboration, which has developed classification criteria for 25 of the most common uveitides using a formal approach to development and classification. Among the diseases studied was MFCPU.^{19–24}

Methods

The SUN Developing Classification Criteria for the Uveitides project proceeded in four phases as previously described: 1) informatics, 2) case collection, 3) case selection, and 4) machine learning.^{21–24}

Informatics.

As previously described, the consensus-based informatics phase permitted the development of a standardized vocabulary and the development of a standardized, menu-driven hierarchical case collection instrument.²¹

Case collection and case selection.

De-identified information was entered into the SUN preliminary database by the 76 contributing investigators for each disease as previously described.^{23,24} Cases in the preliminary database were reviewed by committees of 9 investigators for selection into the final database, using formal consensus techniques described in the accompanying article.^{23,24} Because the goal was to develop classification criteria,²⁵ only cases with a supermajority agreement (>75%) that the case was the disease in question were retained in the final database (i.e. were “selected”).^{23,24}

Machine learning.

The final database then was randomly separated into a training set (~85% of the cases) and a validation set (~15% of the cases) for each disease as described in the accompanying article.²⁴ Machine learning was used on the training set to determine criteria that minimized misclassification. The criteria then were tested on the validation set; for both the training set and the validation set, the misclassification rate was calculated for each disease. The misclassification rate was the proportion of cases classified incorrectly by the machine learning algorithm when compared to the consensus diagnosis. For MF CPU, the diseases against which it was evaluated were: acute posterior multifocal placoid pigment epitheliopathy (APMPPE), BSCR, multiple evanescent white dot syndrome (MEWDS), PIC, serpiginous choroiditis, sarcoidosis-associated posterior uveitis, syphilitic posterior uveitis, and tubercular uveitis.

The study adhered to the principles of the Declaration of Helsinki. Institutional Review Boards (IRBs) at each participating center reviewed and approved the study; the study typically was considered either minimal risk or exempt by the individual IRBs.

Results

Two hundred fifty-one cases of MF CPU were collected, and 138 (57%) achieved supermajority agreement on the diagnosis during the “selection” phase and were used in the machine learning phase. These cases of MF CPU were compared to cases of posterior uveitides, including 82 cases of APMPPE, 207 cases of BSCR, 51 cases of MEWDS, 122 cases of serpiginous choroiditis, 144 cases of PIC, 12 cases of sarcoid posterior uveitis, 35 cases of syphilitic posterior uveitis, and 277 cases of tubercular posterior or panuveitis uveitis. The details of the machine learning results for these diseases are outlined in the accompanying article.²³ The characteristics of cases with MF CPU are listed in Table 1, and the classification criteria developed after machine learning are listed in Table 2. Key features of the criteria include multifocal choroiditis with round or oval lesions >125 µm in size, involvement of the mid-periphery and/or periphery, and punched-out atrophic scars (Figure 1) or active lesions with more than minimal vitritis. The overall accuracies for posterior uveitides were 93.9% in the training set and 98.0% (95% confidence interval 94.3, 99.3) in the validation set. The misclassification rate for MF CPU in the training set was 15%, and in the validation set 0%. The diseases with which MF CPU most often was confused in the training set were BSCR and PIC.

Discussion

The classification criteria developed by the SUN Working Group for MFPCU have a reasonably low misclassification rate, indicating good discriminatory performance against other posterior uveitides.

Multifocal choroiditis is an ambiguous term, which can refer to a clinical finding, a class of diseases, or to a specific disease.²⁶ Some clinicians have used multifocal choroiditis to refer all posterior uveitides with choroidal involvement, others to MFPCU without anterior chamber or vitreous inflammation, and others as synonymous with MFPCU. Although not all cases of MFPCU have anterior chamber or vitreous inflammation, we prefer to use the term MFPCU as the diagnostic entity (regardless of the amount of anterior chamber and vitreous inflammation) and the term “multifocal choroiditis” as a clinical finding indicating multifocal choroidal inflammatory lesions or as a class of diseases characterized by multifocal choroidal inflammation (the multifocal choroiditides). Despite its name, which is used both for historical reasons and to avoid confusion with the use of multifocal choroiditis as a clinical finding or class of diseases, MFPCU is classified as a posterior uveitis, as its primary site of inflammation is in the choroid.^{19,20}

In 1984 Watzke et al²⁷ described the disease known as PIC, characterized by “punctate” choroidal lesions, typically <250 µm in size and often <125 µm in size, no to minimal anterior chamber and vitreous inflammation, and a high rate of choroidal neovascularization.² Because of the rare occurrence of PIC-like lesions in one eye and MFPCU in the other, and because of a similar appearance on multi-modal imaging (other than lesion size), some investigators have considered PIC and MFPCU to be variants of the same disease.^{26,28} Conversely, other investigators, classifying the two diseases based solely on chorioretinal morphology have found clear cut differences, namely the absence of anterior chamber and vitreous inflammation and the absence of uveitis-related structural complications other than choroidal neovascularization in PIC,² and differences in the course with prognostic import.²⁰ There also are differences in the location of the inflammatory lesions/scars; cases of MFPCU typically have mid-peripheral or peripheral involvement, whereas the lesions in PIC typically are concentrated in the posterior pole.^{30,31} A study of cases of MFPCU and PIC using cluster analysis determined that two distinct clusters existed, conforming to the diagnoses of PIC and MFPCU and that the two distinguishing features were anterior chamber and vitreous inflammation (largely absent in PIC) and lesion location (posterior in PIC and peripheral in MFPCU).³⁰

In the series by Shimada et al,³² histological evaluation of surgically-removed choroidal neovascular membranes removed surgically demonstrated inflammatory infiltrates in some cases of MFPCU but not in PIC, suggesting that they might be distinct diseases. Conversely, in the case series by Olsen et al³³ histological evaluation of surgically-removed choroidal neovascular membranes from patients with PIC demonstrated the occasional lymphocyte, suggesting that the pathology may not be completely dissimilar from that of MFPCU. A genetic risk factor association study suggested similar haplotype associations in the *IL-10* and *TNF* loci for MFPCU and PIC, suggesting possible similarities in pathogenesis, but allowing for different inciting events or epigenetic factors to influence phenotype.²⁹ Finally,

if they were a single disease, one might expect the clinical presentation to be a Gaussian distribution with the overlap syndrome to be the most common presentation, which is not the case. The paradigms are the most common presentations, and overlap is uncommon. Hence, the SUN Working Group has elected to define the diseases separately, recognizing that there will be cases with an overlap appearance. Most such cases behave more like MFCPU than PIC and might be classified as MFCPU, but probably should be classified as an overlap syndrome at this time.

The presence of any of the exclusions in Table 2 suggests an alternate diagnosis, and the diagnosis of MFCPU should not be made in their presence. In prospective studies, many of these tests will be performed routinely, and the alternative diagnoses excluded. However, in retrospective studies based on clinical care, not all of these tests may have been performed. In these studies the presence of an exclusionary criterion excludes MFCPU, but the absence of such testing does not always exclude the diagnosis of MFCPU if the criteria for the diagnosis are met

Classification criteria are employed to diagnose individual diseases for research purposes.²⁵ Classification criteria differ from clinical diagnostic criteria, in that although both seek to minimize misclassification, when a trade-off is needed, diagnostic criteria typically emphasize sensitivity, whereas classification criteria emphasize specificity,²⁵ in order to define a homogeneous group of patients for inclusion in research studies and limit the inclusion of patients without the disease in question that might confound the data. The machine learning process employed did not explicitly use sensitivity and specificity; instead it minimized the misclassification rate. Because we were developing classification criteria and because the typical agreement between two uveitis experts on diagnosis is moderate at best,²¹ the selection of cases for the final database (“case selection”) included only cases which achieved supermajority agreement on the diagnosis. As such, some cases which clinicians would diagnose with MFCPU may not be so classified by classification criteria.

In conclusion, the criteria for MFCPU outlined in Table 2 appear to perform sufficiently well for use as classification criteria in clinical research.²⁴

Grant support:

Supported by grant R01 EY026593 from the National Eye Institute, the National Institutes of Health, Bethesda, MD, USA; the David Brown Fund, New York, NY, USA; the Jillian M. And Lawrence A. Neubauer Foundation, New York, NY, USA; and the New York Eye and Ear Foundation, New York, NY, USA.

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Figure 1. Fundus photograph of a case multifocal choroiditis with panuveitis, demonstrating multiple punched-out atrophic chorioretinal lesions.

Table 1.

Characteristics of Cases with Multifocal Choroiditis with Panuveitis

Characteristic	Result
Number cases	138
<i>Demographics</i>	
Age, median, years (25 th 75 th percentile)	38 (28, 55)
Gender (%)	
Men	22
Women	78
Race/ethnicity (%)	
White, non-Hispanic	71
Black, non-Hispanic	9
Hispanic	2
Asian, Pacific Islander	3
Other	4
Missing	11
<i>Uveitis History</i>	
Uveitis course (%)	
Acute, monophasic	3
Acute, recurrent	1
Chronic	92
Indeterminate	4
Laterality (%)	
Unilateral	13
Unilateral, alternating	0
Bilateral	87
<i>Ophthalmic examination</i>	
Keratic precipitates (%)	
None	91
Fine	4
Round	3
Stellate	0
Mutton Fat	1
Other	1
Anterior chamber cells (%)	
Grade 0	54
½+	24
1+	12
2+	6
3+	3

Characteristic	Result
4+	1
Anterior chamber flare (%)	
Grade 0	78
1+	19
2+	3
3+	0
4+	0
Iris (%)	
Normal	96
Posterior synechiae	4
Iris nodules	
Iris atrophy (sectoral, patchy, or diffuse)	0
Heterochromia	0
Intraocular pressure (IOP), involved eyes	
Median, mm Hg (25 th , 75 th percentile)	16 (13, 18)
Proportion patients with IOP>24 mm Hg either eye (%)	4
Vitreous cells (%)	
Grade 0	29
½+	15
1+	36
2+	19
3+	1
4+	0
Vitreous haze (%)	
Grade 0	58
½+	15
1+	16
2+	8
3+	3
4+	0
<i>Chorioretinal lesion characteristics</i>	
Lesion number (%)	
Unifocal (1 lesion)	0
Paucifocal (2–4)	5
Multifocal (5)	89
Missing	6
Lesion shape & character (%)	
Ameboid or serpentine	0
Oval or round	94

Characteristic	Result
Placoid	0
Punched-out/atrophic scars	78
Punctate	0
Missing	6
Inflammatory lesion/scar location (%) [*]	
Posterior pole only involved	1.5
Posterior pole and periphery/mid-periphery	56.5
Mid-periphery and periphery only	42
Typical lesion size (%)	
<125 µm	0
125–250 µm	33
250–500 µm	37
>500 µm	23
Missing	7
Other features (%)	
Peripapillary atrophy	39
Retinal vascular sheathing	9
Retinal vascular leakage	13
Choroidal neovascularization	7

^{*}Based on 129 cases with photographs.

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Table 2.

Classification Criteria for Multifocal Choroiditis with Panuveitis

Criteria
1. Multifocal choroiditis with
a. Oval or round lesions AND
b. Predominant lesion size >125 μ m
AND
2. Characteristic appearance
a. "Punched-out atrophic" chorioretinal scars OR
b. Active lesions with more than minimal vitreous inflammation
AND
3. Inflammatory lesions and/or characteristic scars involving the mid-periphery or periphery with or without posterior pole involvement
Exclusions
1. Positive serologic test for syphilis using a treponemal test
2. Evidence of sarcoidosis (either bilateral hilar adenopathy on chest imaging or tissue biopsy demonstrating non-caseating granulomata)
3. In tuberculosis-endemic regions or tuberculosis-exposed individuals [*] , evidence of infection with <i>Mycobacterium tuberculosis</i>
a. Histologically- or microbiologically-confirmed infection with <i>M. tuberculosis</i> [†] OR
b. Positive interferon- γ release assay (IGRA) [‡] OR
c. Positive tuberculin skin test [§]

^{*} Testing not needed in tuberculosis non-endemic regions.

[†] E.g. biopsy, fluorochrome stain, culture, or polymerase chain reaction based assay.

[‡] E.g. Quantiferon-gold or T-spot.

[§] E.g. purified protein derivative; a positive result should be >10 mm induration