

Mayo Clinic consensus report on membranous nephropathy: proposal for a novel classification



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Membranous nephropathy (MN) is a pattern of injury caused by autoantibodies binding to specific target antigens, with accumulation of immune complexes along the subepithelial region of glomerular basement membranes. The past 20 years have brought revolutionary advances in the understanding of MN, particularly via the discovery of novel target antigens and their respective autoantibodies. These discoveries have challenged the traditional classification of MN into primary and secondary forms. At least 14 target antigens have been identified, accounting for 80%–90% of cases of MN. Many of the forms of MN associated with these novel MN target antigens have distinctive clinical and pathologic phenotypes. The Mayo Clinic consensus report on MN proposes a 2-step classification of MN. The first step, when possible, is identification of the target antigen, based on a multistep algorithm and using a combination of serology, staining of the kidney biopsy tissue by immunofluorescence or immunohistochemistry, and/or mass spectrometry methodology. The second step is the search for a potential underlying disease or associated condition, which is particularly relevant when knowledge of the target antigen is available to direct it. The meeting acknowledges that the resources and equipment required to perform the proposed testing may not be generally available. However, the meeting consensus was that the time has come to adopt an antigen-based classification of MN because this approach will allow for accurate and specific MN diagnosis, with significant implications for patient management and targeted treatment.

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Classifying and ranking is the highest activity the mind can perform.

It is the activity that implies all others.

Paul Otlet¹

Membranous nephropathy (MN) is a common lesion that causes nephrotic syndrome in children and adults. MN is characterized by deposition of immune complexes along the subepithelial region of the glomerular basement membranes (GBMs). A kidney biopsy traditionally has been required to confirm the diagnosis of MN. On light microscopy, the GBM is thickened, and basement membrane spikes or pin holes can be seen along the GBM, although these changes may be absent in early MN. Immunofluorescence (IF) and/or immunohistochemistry (IHC) studies show granular staining for IgG, usually in association with C3 along the capillary walls. Electron microscopy confirms the diagnosis by demonstrating the presence

of subepithelial electron dense deposits. Thus, the hallmark of the MN lesion is the presence of immune complexes, presumably formed *in situ*, composed of IgG and target antigens.

The past 20 years have brought significant and revolutionary advances in the understanding of the nature of MN, particularly through the discovery of target antigens and their respective autoantibodies.^{2–5} In response to these remarkable advances, a group of renal pathologists and nephrologists met at the Mayo Clinic, in Rochester, Minnesota, on October 29, 2022, to reach a consensus on the nomenclature/classification and reporting of MN, and to propose a novel classification. The meeting opened with a review of the historical and clinical perspectives on MN, followed by discussion of antigens, including cell biology and animal models, and the role of complement. The final discussion focused on both the need for and a proposal for standardization of the classification of MN. This report describes the conclusions resulting from this meeting.

THE TRADITIONAL CLASSIFICATION OF MN

MN is not a specific diagnosis *per se*, but rather a pattern of glomerular injury that may be caused by or associated with a variety of disease entities. The common denominator is the deposition of immune complexes along the subepithelial aspect of the GBM. The MN lesion is traditionally classified as being either *primary* MN (~70%) or *secondary* MN (~30%).^{6–9} Primary MN is a classification used when the MN lesion is not associated with any potentially causal and recognizable systemic disease, infection, or drug exposure, and therefore appears as a renal-limited disease.⁶ By contrast, secondary MN is a classification associated with systemic diseases, such as autoimmune diseases, infections, neoplasia, hematopoietic stem cell transplantation, sarcoidosis, and drugs and/or toxins. The relevance of this classification pertains to the treatment of MN. Immunosuppressive treatment is required in many cases of primary MN with nephrotic syndrome, whereas management of secondary MN is based primarily on treatment of the underlying associated disease.

The discovery of the target antigen M-type phospholipase A2 receptor (PLA2R) and its corresponding serum anti-PLA2R antibodies,³ followed by identification of additional novel target antigens in patients with apparent primary MN, has shifted the paradigm of MN classification. The traditional primary versus secondary dichotomy was not changed fundamentally, but rather was refined into a classification more aligned with the evolving understanding of the underlying pathogenesis and diverse number of target antigens. Conceptually, the primary form of MN occurs when circulating antibodies, mostly belonging to the IgG4 subclass, target endogenous antigens expressed on podocytes, although even this simple definition may not hold in all cases. Most cases of primary MN are associated with IgG–PLA2R immune-complex deposits in glomeruli and circulating anti-PLA2R auto-antibodies.^{10–13} In secondary MN, the immune complexes have been hypothesized to consist of a foreign antigen or a neo-epitope that is planted in the subepithelial

space and serves as a target for circulating antibodies. Alternatively, circulating immune complexes theoretically could deposit on the luminal side of the GBM, dissociate, and reform in the subepithelial space, but this possibility has been difficult to demonstrate in human disease. Secondary MN remained essentially a clinical classification, based on identification of a possible underlying disease, until the recent discovery of novel target antigens in an increasing proportion of cases of secondary MN.

THE JOURNEY OF ANTIGEN DISCOVERY

The discovery of PLA2R as a major target antigen in MN in 2009 resulted in a profound change in the approach to MN.³ PLA2R antigen staining in kidney biopsy material and detection of circulating anti-PLA2R antibodies became routine tests, and most cases of primary MN are now classified—based on whether PLA2R is detected in the kidney biopsy specimen and/or circulating anti-PLA2R antibodies—as PLA2R-positive or PLA2R-negative MN.^{10,11,14} PLA2R is detected in approximately 50%–80% of primary MN (depending on age and geography) but also, though rarely, in secondary forms of MN. The remaining approximately 40% of all MN cases (primary and secondary) are PLA2R-negative. Thrombospondin type-I domain-containing 7A (THSD7A) was discovered in 2014, but it is seen in only a small fraction of cases of MN (both primary and associated with neoplasms).⁴ Although many centers perform staining of biopsy tissue for THSD7A, serologic testing is limited to specialized laboratories. Over the past

5 years, many additional, novel target antigens have been discovered,⁵ thereby filling the “antigen void” of PLA2R-negative MN (Figure 1, Table 1). Unlike the initial discovery of anti-PLA2R antibodies in association with MN via traditional biochemical and immunologic approaches,³ most of these additional antigens were identified via laser-capture microdissection of affected glomeruli, in which the target antigen had accumulated within immune deposits, followed by digestion with proteolytic enzymes and analysis of the resulting enriched peptides by tandem mass spectrometry.^{5,15–20} Some of the new target antigens have no known clinical associations and appear to represent primary MN; some are specific to a distinct etiology of secondary MN. Furthermore, different target antigens may be associated with the same secondary disease.²¹ What also became apparent is that many of the MN types associated with novel target antigens have distinctive epidemiologic, clinical (including likelihood of spontaneous remission), and pathologic features and that grouping them together under the generic term “PLA2R-negative MN” would thus be imprecise and uninformative. The discovery of the corresponding serum antibodies to the novel antigens provides important new opportunities for the development of serum tests to facilitate diagnosis and monitoring of the response to treatment.

Antigens most often associated with primary MN

M-type phospholipase A2 receptor (PLA2R). PLA2R was discovered in 2009 and is the most common target in MN.³

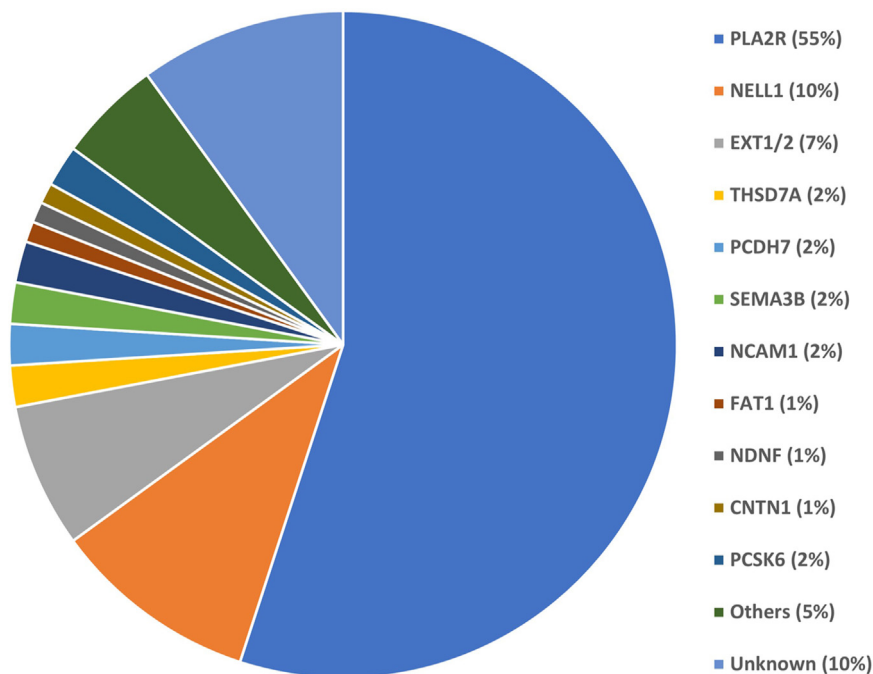


Figure 1 | Approximate incidence of membranous nephropathy (MN) antigens. The target antigen incidence can vary based on demographics, and the incidence (%) shown here is based on cases of M-type phospholipase A2 receptor (PLA2R)–negative MN used for mass spectrometry detection of MN antigens at the Mayo Clinic, Rochester, Minnesota, USA. Others include rare (< 1%) and putative antigens. Neither serine protease HTRA1 nor transforming growth factor beta receptor 3 were detected in any case, on mass spectrometry studies, and hence, they are not shown. CNTN1, contactin 1; EXT, exostosin; FAT1, protocadherin FAT1; NCAM1, neural cell-adhesion molecule 1; NDNF, neuron-derived neurotrophic factor; NELL1, neural epidermal growth factor–like protein 1; PCDH7, protocadherin 7; PCSK6, proprotein convertase subtilisin/kexin type 6; SEMA3B, semaphorin 3B; THSD7A, thrombospondin type-I domain-containing 7A.

Table 1 | Characteristics of (known) MN target antigens

Target antigen	Podocyte expressed? TM vs. secreted	AutoAb detected?	Clinical/disease association	Distinctive histopathologic features
PLA2R	Yes; TM	Yes	None	Global, granular, subepithelial deposits; IgG4 predominant
THSD7A	Yes; TM	Yes	Malignancy	Similar to PLA2R Caveat: there is normal linear basal podocyte staining for THSD7A
NELL1	No ^a ; secreted	Yes	Malignancy, drugs, autoimmune	IgG1 predominant, deposits may be segmental or incomplete-loop pattern
SEMA3B	Yes; secreted	Yes	Pediatric	IgG1 predominant, may have additional mesangial deposits; TBM deposits may be present
PCDH7	No; TM	Yes	Older	C3 absent or weak
HTRA1	Yes; secreted	Yes	None	IgG4 predominant, similar to PLA2R
NTNG1	Yes; GPI-linked	Yes	None	IgG4 predominant, similar to PLA2R
EXT1/EXT2	No ^a ; TM Golgi protein and secreted	No	Autoimmune disease, lupus	IgG1 predominant, IgA, IgM often present, mesangial deposits, may coexist with class III/IV lupus
NCAM1	No ^a ; TM	Yes	Lupus	Similar to EXT1/EXT2
TGFBR3	Yes; TM	No ^b	Lupus	Similar to EXT1/EXT2
CNTN1	No ^a ; GPI-linked	Yes	CIDP	IgG4 predominant
FAT1	Yes; TM	Yes	HSCT	TBM deposits can be present
NDNF	Yes; secreted	Yes	Young males, syphilis	Lumpy deposits, superficial hump-like by EM, IgG1
PCSK6	No ^a ; secreted	Yes	Prolonged NSAID use	IgG1 and 4

AB, antibodies; CIDP, chronic inflammatory demyelinating polyradiculoneuropathy; CNTN1, contactin 1; EM, electron microscopy; EXT, exostosin; FAT1, protocadherin FAT1; GPI, glycosylphosphatidylinositol (links protein to membrane); HSCT, hematopoietic stem cell transplant; HTRA1, serine protease HTRA1; MN, membranous nephropathy; NCAM1, neural cell-adhesion molecule 1; NDNF, neuron-derived neurotrophic factor; NELL1, neural epidermal growth factor-like protein 1; NSAID, nonsteroidal anti-inflammatory drug; NTNG, netrin G1; PCDH7, protocadherin 7; PCSK6, proprotein convertase subtilisin/kexin type 6; PLA2R, M-type phospholipase A2 receptor; SEMA3B, semaphorin 3B; TBM, tubular basement membrane; TGFBR3, transforming growth factor beta receptor 3; THSD7A, thrombospondin type-I domain-containing 7A; TM, transmembrane.

^aThese antigens are not well-expressed by the normal podocyte according to current single-cell transcriptomic data sets, although it is possible that they are upregulated in disease.

^bAntigen was identified by tissue immunostaining (bound to IgG).

PLA2R-associated MN is characterized by granular staining for PLA2R along the GBM, anti-PLA2R antibodies predominantly of the IgG4 subclass (in most cases), and detection of anti-PLA2R antibodies in the serum. Important to note is that the disease activity correlates over time with the titer and trajectory of antibodies, and the titers decline with successful immunosuppression.²² In a consecutive series of adult patients with biopsy-proven MN with available identification of the target antigen, as well as conditions traditionally associated with MN, 81% of patients were classified as having PLA2R-associated MN. The majority of these patients had no associated disease and behaved as would a typical patient with primary MN.¹³

Thrombospondin type-1 domain-containing 7A (THSD7A).

THSD7A-associated MN was discovered in 2014, and in the majority of cases, it is a prototypical example of a primary MN, with anti-THSD7A antibodies, predominantly of the IgG4 subclass (in most cases), directed against THSD7A that is endogenously expressed on podocytes.⁴ THSD7A accounts for approximately 1%–3% of cases.²³ As with PLA2R, the disease activity correlates with the titer and trajectory of antibodies and responds to immunosuppression. An important finding is that a substantial minority of patients with this form of MN have a concurrent malignancy; a large series showed malignancy in 31 of 195 patients (16%) with THSD7A-associated MN.²⁴ Although only a minority of patients had a clear temporal association between malignancy and the diagnosis of MN, in a few cases, the causal relationship was unequivocal.²⁵ Circulating anti-THSD7A antibodies have been reported as a consequence of THSD7A expression by a malignant tumor.

These cases therefore do not fit well within the current primary and secondary MN classifications.

Neural epidermal growth factor-like protein 1 (NELL1).

NELL1 was identified as an MN target antigen in 2019.¹⁶ NELL1-associated MN is characterized by NELL1 staining along the GBM; in a subset, only segmental GBM staining is present.²⁶ The anti-NELL1 antibody belongs to primarily the IgG1 subclass, and anti-NELL1 antibodies are detectable in the serum. NELL1 appears to be the second-most common antigen in MN, accounting for approximately 10% of cases. An interesting finding is that although most cases of NELL1-associated MN do not have an underlying associated disease, NELL1 has been identified in many secondary forms of MN.^{27,28} Depending on the case series, 10%–33% of NELL1-associated MN is associated with malignancy.²⁷ NELL1-associated MN also has been reported to occur with alpha-lipoic acid use and after mercury exposure due to use of traditional indigenous medicines from India.^{29,30} Important to note is that many patients achieve remission after cessation of use of the lipoic acid or traditional medicine.

Semaphorin 3B (SEMA3B). SEMA3B was identified as an MN target antigen in 2020.¹⁷ SEMA3B-associated MN is unique in that it appears to be present in mostly the pediatric age group. It is characterized by SEMA3B staining along the GBM; the anti-SEMA3B antibody belongs to primarily the IgG1 subclass; and anti-SEMA3B antibodies are detectable in the serum. Unlike antibodies identified in other forms of MN, the anti-SEMA3B antibodies react with only the reduced antigen, thus suggesting that they recognize a cryptic epitope. Tubular

basement membrane immune deposits may be present, but they are negative for SEMA3B staining. SEMA3B accounts for approximately 2% of MN, but it appears to have a higher incidence in pediatric MN. Recurrent SEMA3B-associated MN in kidney allografts has been reported recently.³¹

Protocadherin 7 (PCDH7). PCDH7 was identified as an MN target antigen in 2021.¹⁹ PCDH7-associated MN appears to be present in the older age group and is characterized by PCDH7 staining along the GBM. The anti-PCDH7 IgG subclass may be either IgG1 or IgG4, and anti-PCDH7 antibodies are detectable in the serum. Interestingly, kidney biopsy shows absent or minimal complement deposition. PCDH7-associated MN likely accounts for approximately 2% of all MN cases.

Serine protease HTRA1 (HTRA1). HTRA1 was discovered in 2021.³² HTRA1-associated MN does not appear to be associated with any disease entity and is characterized by HTRA1 staining along the GBM. The anti-HTRA1 IgG subclass is IgG4, and anti-HTRA1 antibodies are detectable in the serum.

Netrin G1 (NTNG1). NTNG1 is a rare MN target antigen discovered in 2022.³³ MN associated with NTNG1 does not appear to be associated with any disease entity and is characterized by NTNG1 staining along the GBM. The anti-NTNG1 IgG subclass is IgG4, and anti-NTNG1 antibodies are detectable in the serum. NTNG1-associated MN likely accounts for less than 1% of all cases.

Antigens most often associated with secondary MN

Exostosin 1 and exostosin 2 (EXT1/EXT2). EXT1/EXT2 were discovered in 2019 as putative antigens in MN associated with autoimmune disease, such as lupus.¹⁵ Circulating antibodies to EXT1/EXT2 have not yet been reported. EXT1/EXT2-associated MN is characterized by staining for both EXT1 and EXT2 along the GBM; the IgG subclass is IgG1; and features of associated autoimmune disease, such as mesangial deposits and endothelial tubuloreticular inclusions, are usually present. EXT1/EXT2 exists as a heterodimer, and IHC/IF demonstrates the presence of both EXT1 and EXT2 in all cases. EXT1/EXT2 likely accounts for 5%–10% of MN cases, and approximately 33%–46% of cases of membranous lupus nephritis are associated with EXT1/EXT2.^{34–36} In addition, EXT1/EXT2-associated MN may coexist with proliferative class III and class IV lupus nephritis. EXT1/EXT2-associated MN is associated with good clinical outcomes, and minimal chronicity on kidney biopsy.^{34–36}

Neural cell-adhesion molecule 1 (NCAM1). NCAM1 was discovered in 2021.³⁷ NCAM1-associated MN appears to be present in a subset of patients with membranous lupus nephritis and in patients with no associated disease. It is characterized by NCAM1 staining along the GBM. The anti-NCAM1 IgG subclass is variable, and anti-NCAM1 antibodies are detected in the serum. Like EXT1/EXT2, NCAM1 may present together with proliferative lupus nephritis classes III and IV. NCAM1-associated MN likely accounts for 1%–2% of all MN cases, and approximately 6% of membranous lupus nephritis cases.³⁷

Transforming growth factor beta receptor 3 (TGFBR3).

TGFBR3 was discovered in 2021 and was detected in 6% of patients with membranous lupus nephritis.³⁸ Granular GBM staining for TGFBR3 was present, but no circulating antibodies were detected.

Protocadherin FAT1 (FAT1). FAT1 was discovered in 2022 and is associated specifically with MN in patients who have received an allogeneic hematopoietic stem cell transplant.¹⁸ FAT1-associated MN is characterized by staining for FAT1 along the GBM; the anti-FAT1 antibody belongs primarily to the IgG4 subclass; and anti-FAT1 antibodies are detected in both the serum and kidney biopsy tissue. As with PCDH7, which is also a cadherin, either minimal or no glomerular staining occurs for complement proteins by IF. Additionally, both PCDH7 and FAT1 are heavily glycosylated, requiring protease pretreatment for IHC/IF demonstration of PCDH7 and FAT1 staining.

Contactin 1 (CNTN1). CNTN1 was discovered in 2020 and appears to be specific for MN associated with chronic inflammatory demyelinating polyneuropathy.^{39–42} CNTN1-associated MN is characterized by staining for CNTN1 along the GBM; the anti-CNTN1 antibody primarily belongs to the IgG4 subclass; and anti-CNTN1 antibodies are detected in both the serum and kidney biopsy tissue.

Neuron-derived neurotrophic factor (NDNF). NDNF was discovered in 2023 and is unique in that it is present in MN patients with syphilis.²⁰ NDNF-associated MN is characterized by staining for NDNF along the GBM; the anti-NDNF antibody belongs primarily to the IgG1 subclass; electron microscopy shows superficial sub-epithelial scattering of deposits (stage I MN); and anti-NDNF antibodies are detected in kidney biopsy tissue (serum was not available for this study). Important to note is that most cases of NDNF-associated MN respond to penicillin treatment of syphilis without the need for immunosuppression.

Proprotein convertase subtilisin/kexin type 6 (PCSK6). PCSK6 was discovered in 2023 and is unique in that it is present mostly in MN patients with prolonged nonsteroidal anti-inflammatory drug use.⁴³ PCSK6-associated MN is characterized by staining for PCSK6 along the GBM; the anti-PCSK6 antibody belongs to both the IgG1 and IgG4 subclasses; and anti-PCSK6 antibodies are detected in kidney biopsy tissue. Important to note is that most cases of PCSK6-associated MN respond to discontinuation of nonsteroidal anti-inflammatory drug use and conservative management.

Other putative antigens. Additional rare (1%–2% each) putative antigens and other candidate proteins have been identified.^{23,44–46} However, confirmatory studies including detailed clinicopathologic findings, validation studies, and demonstration of corresponding serum antibodies remain to be done in many of these candidate proteins.

For most new antigens, circulating antibodies have been found that have implicated the proteins as true target antigens. For others, antibodies have not yet been found, which

could be due to either technical and methodological reasons or the fact that a particular protein accumulates for other reasons, such as being part of the pathologic process but not the target antigen *per se*. In this report, we have used the term “target antigen” loosely, with the understanding that the roles of these proteins in the disease process will be better elucidated with time.

THE NEED FOR A NEW CLASSIFICATION OF MN

A classification must reflect the underlying pathophysiology of the disease, but at the same time, it must be pragmatic and guide clinicians in their diagnostic, prognostic, and therapeutic approaches. Over the past few years, the portfolio of target antigens has broadened substantially. Certain antigens cannot be assigned as being exclusively representative of primary or secondary disease. Conversely, other target antigens appear to occur in the context of a specific exposure, associated conditions, or demographic profile. As the traditional primary–secondary MN dichotomy loses its relevance, we propose a novel classification that is based on the target antigen and associated disease.

Although substantial advances have occurred in the identification of antigens in childhood MN, they were not specifically discussed during this meeting.⁷ However, the need for a new classification and the proposed terminology applies similarly to both pediatric and adult MN.

A PROPOSAL FOR AN ANTIGEN- AND DISEASE-BASED CLASSIFICATION (2-STEP)

Step 1: antigen association

We propose that efforts should be made to identify the associated target antigen in each case of MN (Figure 2). The underlying target antigen can be determined by tandem mass spectroscopy in approximately 80%–90% of MN cases. IF/IHC are now in widespread use for identification of PLA2R and THSD7A, and in increasing use for identification of NELL1. As additional antibodies become available and are validated, an immunohistochemical approach can be advocated increasingly, with a logical cascade, starting with PLA2R and possibly 1 or 2 others, followed by a more directed approach if tests for these are negative.

Serology also can be used to determine the target antigen by demonstrating the presence of the corresponding antibody.¹² Thus, identification of serum anti-PLA2R or anti-THSD7A antibodies would be consistent with PLA2R- and THSD7A-associated MN. Furthermore, among patients with nephrotic syndrome and (i) preserved kidney function and (ii) no evidence of secondary causes, including diabetes mellitus, a positive PLA2R-antibody test is highly predictive of a tissue diagnosis of PLA2R-associated MN, and a biopsy/tissue-based diagnosis of PLA2R-associated MN is not required.^{10,11,47}

In cases for which a specific target antigen can be established, we recommend that the following terminology be used: MN, [target antigen]–associated. Examples are:

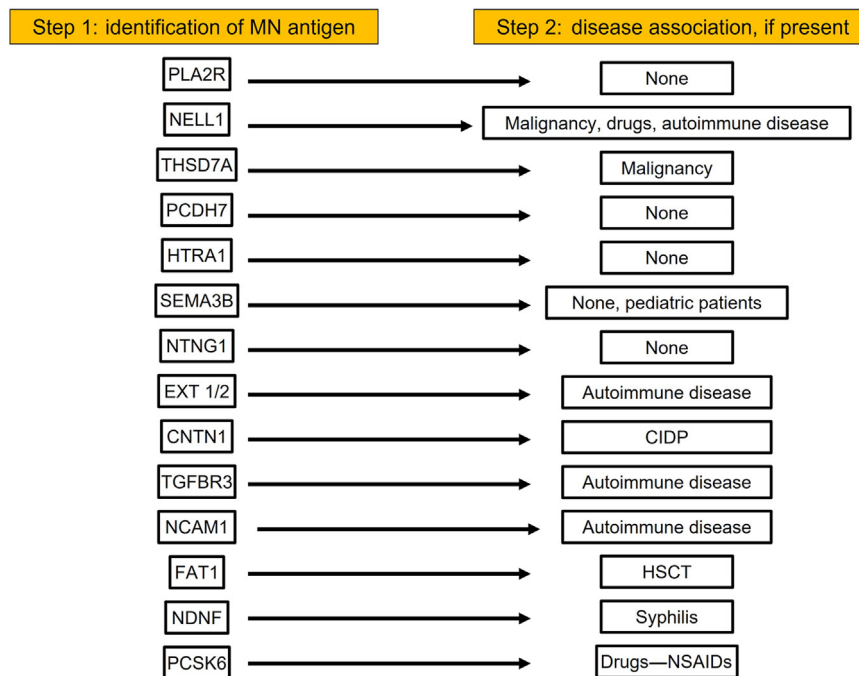


Figure 2 | Proposal for 2-step classification of membranous nephropathy (MN). Antigen/antibody detection in MN is the first step. Identification and correlation with an associated disease or condition, if present, is the second step. CIDP, chronic inflammatory demyelinating polyradiculoneuropathy; CNTN1, contactin 1; EXT, exostosin; FAT1, protocadherin FAT1; HSCT, hematopoietic stem cell transplant; HTRA1, serine protease HTRA1; NCAM1, neural cell-adhesion molecule 1; NDNF, neuron-derived neurotrophic factor; NELL1, neural epidermal growth factor–like protein 1; NSAID, nonsteroidal anti-inflammatory drug; NTNG1, netrin G1; PCDH7, protocadherin 7; PCSK6, proprotein convertase subtilisin/kexin type 6; PLA2R, M-type phospholipase A2 receptor; SEMA3B, semaphorin 3B; TGFBR3, transforming growth factor beta receptor 3; THSD7A, thrombospondin type-I domain-containing 7A.

“MN, PLA2R-associated”; “MN, NELL1-associated”; and “MN, EXT1/EXT2-associated.” In cases in which careful evaluation of an MN lesion does not reveal a known antigen, we suggest using the following terminology: *MN, antigen undetermined*. In cases in which an evaluation for associated antigens was not performed, we suggest using the following terminology: *MN, antigen evaluation not performed*. In cases in which evaluation of antigens was done for a limited number of antigens, all of which were negative, we suggest using the following terminology: *MN, [and list the antigens tested that are negative]*—e.g., “MN, PLA2R/NELL1/EXT-negative,” with a comment that only these specific antigens were tested.

Step 2: disease association

As discussed above, many of the novel antigens have been associated with specific diseases or exposures. These include EXT1/EXT2, NCAM1, FAT1, CNTN1, NDNF, and PCSK6. NELL1 and THSD7A may be present either with or without disease association.

Thus, we propose that if a concurrent disease or exposure appears to be associated with the MN antigen, it should be specified, as in the following examples: MN, PLA2R-associated (no known disease association); MN, PCDH7-associated (no known disease association); MN, THSD7A-associated (prostate carcinoma/clinical); MN, NELL1-associated (lung adenocarcinoma/clinical); MN, EXT1/EXT2-associated (lupus/clinical); MN, NCAM1-associated (lupus/clinical); MN, NDNF-associated (syphilis/clinical); MN, FAT1-associated (hematopoietic stem cell transplant/clinical); MN, CNTN1-associated (polyneuropathy/clinical); and MN, PCSK6-associated (nonsteroidal anti-inflammatory drug use/clinical; [Figure 2](#)).

Given the rarity of certain antigens, evaluation for possible associated conditions can be suggested if it has not been done already. Thus, evaluation for malignancy should be suggested in patients that have NELL1- or THSD7A-associated MN;

evaluation for autoimmune disease should be suggested in patients with EXT1/EXT2- or NCAM1-associated MN, etc.

Disease association is an important component of the traditional classification of MN, and step 2 aims to retain in principle the primary and secondary forms of MN, with correlation of specific antigens associated with specific diseases. As the identification of many antigens is recent, the number of patients with the relevant form of MN is small, and future studies of larger cohorts of patients with a longer follow-up are required. Further development and adjustment of this evolving classification, as new knowledge becomes available, will be important. Age-appropriate malignancy evaluation should also be performed for MN that is not associated with NELL-1 and THSD7A.

TERMINOLOGY

The pathogenicity of antibodies against the novel target antigens has not yet been studied in detail (except for PLA2R, THSD7A, and SEMA3B), and future studies are likely to show that pathogenic mechanisms differ for different antigens. Thus, at the present time, the best approach is to use the term “associated” with each of the antigens.

CLINICAL IMPLICATIONS

Diagnosis and the ultimate classification of disease require the coalescence of information from many sources ([Figure 3](#)). Histopathology of the kidney biopsy has been the primary method of diagnosis for MN, and features to suggest primary disease versus the possibility of associated diseases remain important. In addition to the traditional histopathologic features, we now can determine, in many cases, the target antigen enriched in the deposits by IF/IHC, which gives rise to the classification scheme described herein. Also, the powerful technique of laser-capture microdissection and mass spectrometric analysis of the tissue can be used to determine uniquely enriched peptides that may detect the known forms of MN or suggest new candidates in the remaining undefined

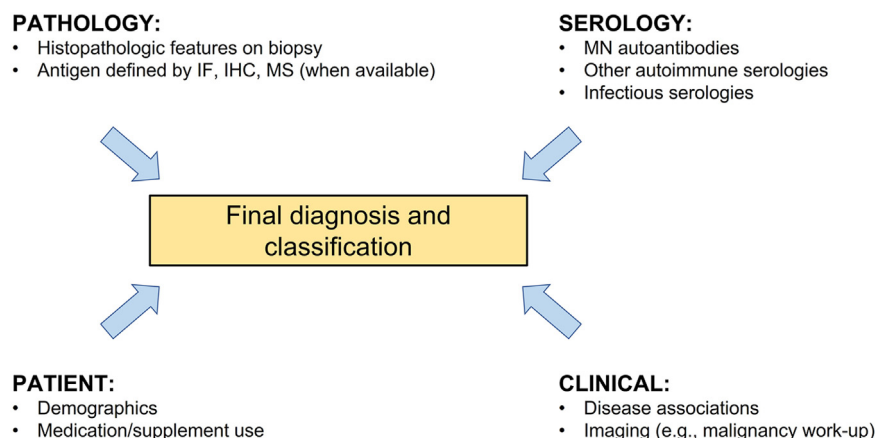


Figure 3 | Integration of data from multiple sources leads to the diagnosis and ultimate classification of membranous nephropathy (MN). Although traditional histopathology of the kidney biopsy and newer techniques to characterize target antigens remain critical for diagnosis, multiple streams of information play a role in identifying circulating autoantibodies and obtaining evidence for associated diseases or medication/supplement use. IF, immunofluorescence; IHC, immunohistochemistry; MS, mass spectrometry.

types of MN. Although the availability, required expertise, and cost associated with mass spectrometry currently may make it available in only select locations, we anticipate wider use of this technology in the near future.

The renal pathologist provides these critical tissue-based data, but other sources of information are needed also. A suggested approach is that, in certain situations, serology can be used to diagnose MN (see next paragraph), obviating the need for biopsy.^{10,11} Thus, our classification system needs to incorporate the emerging serologic tests, such as the test for presence of anti-PLA2R antibodies. This system also incorporates clinical information about associated diseases and medication use that may suggest a secondary association. Information from the patient, primary care physician, and nephrologist must be available for this to occur, and thus, the final diagnosis as listed will need to evolve as this information becomes available to the pathologist and pending studies and queries recommended based on biopsy findings.

In addition to suggesting specific known or unknown secondary disease associations that may warrant treatment or further investigation, determination of the target antigen has important clinical implications. For example, discovery of PLA2R and serum anti-PLA2R auto-antibodies has changed, and continues to change, how we diagnose and treat PLA2R-associated MN.^{10,11,22,48–50} Similar studies subsequently were performed on THSD7A and corresponding anti-THSD7A autoantibodies.^{24,51,52} We anticipate that the detection of new antigens and the corresponding antibodies in serum also will prove critical for diagnosis, treatment, and follow-up. The new antigens will be associated with only rare, and at times unique, subsets of MN cases, emphasizing the need for collaborative efforts in detection of new antigens and corresponding antibodies, treatment, and clinical outcomes.

EVALUATION OF MN ANTIGENS

Testing to detect PLA2R by IF/IHC in kidney biopsy tissue is now routinely performed. Tests to detect some of the newer antigens, such as THSD7A, NELL1, and EXT1/EXT2 also is performed by many laboratories. However, cost and/or lack of availability of IF/IHC for the new antigens can be a major limiting factor for many institutions, even though testing for these antigens is likely to increase in the coming years. Based on the institution and resources, a flexible approach can be adapted, to accommodate the accessibility of antigen testing. The 3 approaches are shown in Figure 4 and are as follows:

Start with basic PLA2R and NELL1 IHC/IF staining, as these are the 2 most common antigens. If negative:

- (1) *Targeted approach*: Perform antigen testing based on clinical disease association. Thus, perform staining studies as follows: for EXT1/EXT2 and NCAM1 if an autoimmune disease is suspected; for FAT1 in patients with HSCT; for CNTN1 in patients with chronic polyneuropathy; for NDNF in patients with syphilis; and for PCSK6 in patients with nonsteroidal anti-inflammatory drug use. In pediatric MN patients, staining for SEMA3B should be performed. If these studies are

negative, or if no established disease association is present, then staining for less-common antigens, including THSD7A, PCDH7, and HTRA1, can be performed. Many of these tests may be sent out to specialized laboratories.

- (2) *Panel approach*: Alternatively, a laboratory could employ a standard panel of antibodies for antigens that can vary depending on volume and demographics, including antigens such as THSD7A, SEMA3B, PCDH7, EXT1/EXT2, NCAM1, FAT1, and PCSK6. If employing IHC or IF staining studies on formalin-fixed paraffin-embedded material, an extra 6–8 slides should be cut upfront for both approaches 1 and 2 to save time and biopsy tissue.
- (3) *Proteomic approach*: Lastly, laser microdissection of MN glomeruli, followed by mass spectrometry, soon will become more widely available as a clinical laboratory test for antigen detection. Data from mass spectrometry also has shown that the antigens are mutually exclusive, and multiple antigens have not been detected in any given case of MN.

An important point to remember is that staining for PLA2R and NELL1 will eliminate the need for the above approach in approximately 60% of cases, and even after the above exhaustive approach is taken, the antigen will remain unknown in approximately 10% of cases. These considerations are important for countries with lower income, and efforts must be made by the community to make the latest techniques available to most centers.

THE KIDNEY BIOPSY REPORT

Briefly, the following should be addressed in the kidney biopsy report when MN is encountered:

- (1) *Pattern-of-injury*: MN. Additional comments on morphologic features suggesting a specific antigen or underlying disease may be included (e.g., segmental lesions, mesangial, subendothelial and/or extra-glomerular deposits, full-house Ig, complement deposition, presence or absence of endothelial tubuloreticular inclusions).
- (2) The target antigen detected; if this is not known, the staining studies that were performed and returned with a negative result should be reported.
- (3) Clinical association and prognostic implications of the target antigens; if this is not known, suggest evaluation in the comment section, depending on the antigen identified.
- (4) IgG subclass description, if performed. IgG subtype can be an ancillary test in confirming an antigen or can guide testing for certain antigens.
- (5) Electron microscopy stage based on Ehrenreich and Churg staging,⁵³ despite the limitations of the staging system.
- (6) Findings that have prognostic implications, including but not limited to secondary focal segmental glomerular sclerosis, extent of glomerulosclerosis, interstitial fibrosis/tubular atrophy, and vascular disease.

Although an individual pathology report is not expected to be a living document that is routinely updated, in some cases we can expect an iterative refinement of the diagnosis

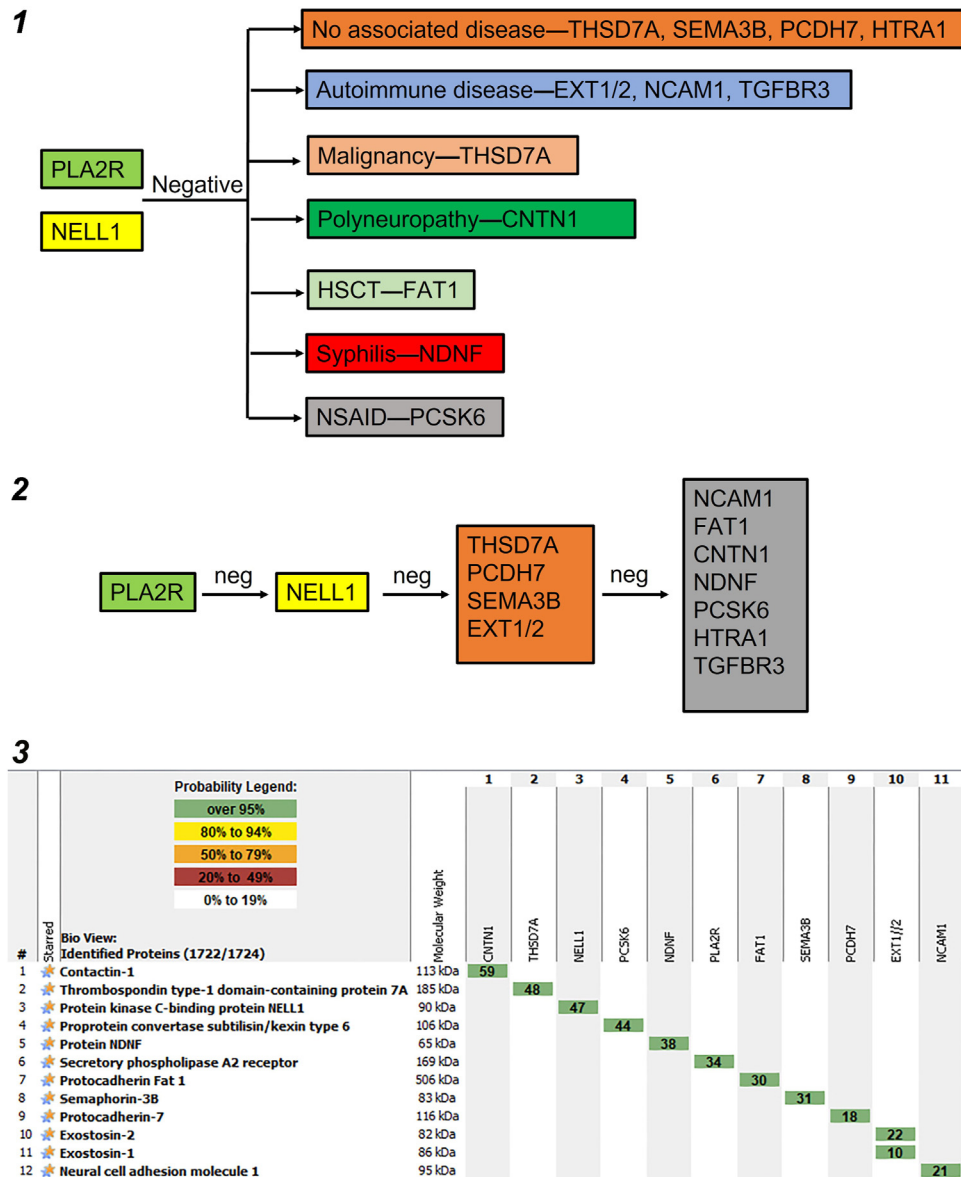


Figure 4 | Approach to detection of target antigen. Initial testing by immunofluorescence/immunohistochemistry (IF/IHC) should be performed for either M-type phospholipase A2 receptor (PLA2R), followed by neural epidermal growth factor–like protein 1 (NELL1), or both together. A positive test for PLA2R or NELL1 indicates membranous nephropathy (MN) associated with PLA2R or NELL1. If both tests are negative, detection of the remaining MN antigens can be done using any of the following: (1) Targeted approach using IF/IHC to screen for specific antigens based on known disease association(s) or absence of any associated disease. (2) Panel approach using IF/IHC of more-common antigens followed by less-common antigens. (3) Proteomic approach using laser microdissection and mass spectrometry. The figure shows total spectral counts (TSCs) of the target antigen in a representative case of MN. Each column represents 1 case of MN; 11 cases are shown side by side. Note that exostosin (EXT)1 and EXT2 are present in the same case. The test should be able to identify a target antigen in greater than 80% of MN cases. The probability number (>95% is highlighted by green) indicates essentially the percent homology between peptides detected in the specimens and the published amino acid sequences of their corresponding proteins. CNTN1, contactin 1; FAT1, protocadherin FAT1; HSCT, hematopoietic stem cell transplant; HTRA1, serine protease HTRA1; NCAM1, neural cell-adhesion molecule 1; NDNF, neuron-derived neurotrophic factor; neg, negative; NSAID, nonsteroidal anti-inflammatory drug; PCDH7, protocadherin 7; PCSK6, proprotein convertase subtilisin/kexin type 6; SEMA3B, semaphorin 3B; TGFBR3, transforming growth factor beta receptor 3; THSD7A, thrombospondin type-I domain-containing 7A.

and classification, with further input from additional pathologic antigen testing (including the anticipated discovery of future target antigens) and/or serologic evaluation, newly recognized clinical associations, testing for infectious or malignant processes, and further discussions with patients and clinicians (Figure 3).

SUMMARY

Significant advances have been made in the understanding, diagnosis, and treatment of MN. These advances have been aided substantially by the discovery of novel antigens in MN, and their importance will increase further as the proposed new classification based on target antigens in MN is

implemented. Once this routine is established more widely, the role of new antigens in MN disease pathogenesis and their associated clinical determinants will rapidly expand and will eventually allow more precision medicine to be used in MN disease management. *With these new advances, classification of MN as primary versus secondary is no longer sufficient; instead, identification and classification of each form of MN, based on the target antigen, are critical.* This consensus report sets forth a proposal on classification that is based on target-antigen identification, along with relevant disease and medication associations, when present.

DISCLOSURE

All the authors declared no competing interests.

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