Clinical Insights into Diffuse Alveolar Hemorrhage in Antiphospholipid Syndrome

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Abstract
Purpose of Review Diffuse alveolar hemorrhage (DAH) is a rare but devastating manifestation of antiphospholipid syndrome (APS) patients with or without other systemic autoimmune diseases. Data regarding diagnosis and treatment are limited to case series. We review diagnostic and therapeutic strategies employed in APS patients with DAH and discuss our experience in managing these complex patients.

Recent Findings Pulmonary capillaritis likely contributes to the pathogenesis, however is only observed in half of the biopsies. Corticosteroids induce remission in the majority of patients, however almost half recur and require a steroid-sparing immunosuppressive to maintain remission. Cyclophosphamide- or rituximab-based regimens achieve the highest remission rates (50%); other strategies include intravenous immunoglobulin, plasmapheresis, mycophenolate mofetil, and/or azathioprine.

Summary Given the rarity of DAH in APS, treatment is guided by interdisciplinary experience. Why certain patients achieve full remission with corticosteroids while others require immunosuppressive agents is unknown; future research should focus on the pathophysiology and optimal management.

Keywords Antiphospholipid syndrome · Antiphospholipid antibodies · Alveolar hemorrhage · Pulmonary hemorrhage · Bleeding · Capillaritis

Introduction
Antiphospholipid syndrome (APS) is a systemic autoimmune disease characterized by thrombosis and obstetric morbidity in patients with persistently elevated antiphospholipid antibodies (aPL) (lupus anticoagulant test [LA], antiphospholipid antibodies [aCL], or anti-β2-glycoprotein-I antibodies [aβ2GPI]) [1]. Thrombosis in APS can be grouped as venous, arterial, or microvascular. Diffuse alveolar hemorrhage (DAH) is a rare and potentially fatal microvascular manifestation of aPL in which red blood cells (RBCs) leak from the alveolar capillaries into the intra-alveolar space [2••]. Patients present with diffuse pulmonary infiltrates, hypoxemic respiratory failure, hemoptyis, and anemia.

Diffuse alveolar hemorrhage in APS is a major therapeutic challenge, particularly in this patient population, who are hypercoagulable and often on life-long anticoagulation. Thus, the goal of this review is to offer insights into the diagnosis as well as the management of aPL-positive patients with DAH.

General Considerations
How Common Is Diffuse Alveolar Hemorrhage in Antiphospholipid Syndrome?

Diffuse alveolar hemorrhage in APS is rare. Hillerdal et al. reported the first patient in 1991 and since then, approximately 100 cases have been published [3]. In a cohort of 1000 APS patients (36% with systemic lupus erythematosus [SLE]), pulmonary hemorrhage, adult respiratory distress syndrome...
(ARDS), and pulmonary artery thrombosis had a combined prevalence of only 0.7% [4]. One institution reported a 2% frequency of aPL-associated DAH among 483 primary APS patients followed over 16 years [5••]. In catastrophic APS (CAPS), DAH is identified in 12% of patients [6]. The nonspecific clinical and radiographic findings and inconsistent hemoptysis in DAH, as well as the requirement for an invasive diagnostic procedure, i.e., bronchoscopy, contribute to DAH being underdiagnosed [7]. Thus, the true frequency is likely higher than what is reported in the literature [8, 9].

What Is the Etiopathogenesis of Diffuse Alveolar Hemorrhage in Antiphospholipid Syndrome?

The defining feature of DAH is injury to the lung’s microcirculation: the capillaries, arterioles, and venules lining the alveoli. Three major histologic descriptions of DAH, not specific for APS, are (1) pulmonary capillaritis, (2) bland alveolar hemorrhage, and (3) diffuse alveolar damage [10••].

Pulmonary capillaritis is defined by neutrophilic infiltration of the perivascular interstitium (alveolar septae) of the capillaries. A vicious cycle of neutrophilic apoptosis, proteolytic enzyme disbursement, fibrin deposition, and recruitment of inflammatory cells culminates in fibrinoid necrosis of the alveolar and capillary walls. Once a critical mass of neutrophils and aforementioned debris disrupt the alveolar-capillary basement membrane, RBCs extravagate into the alveoli and interstitium. Hemosiderin-filled macrophages begin to accumulate 24–48 h after the initial vessel injury [10••, 11, 12]. Bland alveolar hemorrhage is characterized by intra-alveolar RBCs without adjacent capillary inflammation or destruction. Diffuse alveolar damage, the primary lesion in ARDS, is defined by intra-alveolar edema and hyaline membranes filled with debris lining the alveoli with sparse inflammation [10••, 13, 14].

The specific mechanism behind DAH in APS is not well defined. The most frequently proposed theory is antibody-mediated pulmonary capillaritis, possibly caused by aPL-induced endothelial cell activation and upregulation of adhesion molecules [15••]. Another postulated mechanism is in situ microvascular thrombosis generating alveoli hemorrhage [16]. The detailed discussion of mechanisms of aPL-mediated DAH and micro-thrombosis can be found elsewhere [7, 15••, 17]; however, recent data suggest (a) aPL activation of the mammalian target of rapamycin (mTOR) kinase leads to endothelium proliferation and consequent vasculopathy, which is evident in aPL-associated nephropathy, and (b) complement activation is an important contributor to aPL-mediated thrombosis; however, complement deposition is rarely seen on lung biopsy [18–20].

How Do Patients with Antiphospholipid Syndrome and Diffuse Alveolar Hemorrhage Present?

Patients with DAH present with hypoxemia, hemoptysis, pleuritic chest pain, and cough. Dyspnea is the most common symptom, and fever may be present. Severity ranges from asymptomatic radiographic abnormalities to respiratory failure. Hemoptysis may be initially absent in a third of patients, because the total alveolar capacity can hold a significant pool of blood in distal airways [10••]. Acute respiratory distress syndrome may develop but is more common in CAPS [21–23].

Recurrent bleeding episodes can lead to mild to moderate (rarely severe) interstitial fibrosis and restrictive lung disease [3, 24, 25]. Development of lung hyperinflation and airflow obstruction is rare but has been reported [3, 26]. Although pulmonary hypertension in APS is primarily a result of chronic pulmonary thromboembolism and vasculopathy, DAH without embolic disease can also elicit pulmonary hypertension [3, 21, 25].

Laboratory findings often demonstrate anemia, leukocytosis, and elevated erythrocyte sedimentation rate as well as C-reactive protein; however, the absence of these findings does not exclude DAH. Imaging may show diffuse or patchy alveolar opacities. Bilateral ground glass opacities on chest computed tomography (CT) are classically seen.

Diagnosis and Evaluation of Patients Presenting With Diffuse Alveolar Hemorrhage

The differential diagnosis for DAH includes infections, medications, systemic vasculitis, heart failure, uremia, and coagulopathies; a detailed review of differential diagnosis can be found elsewhere [10••].

Infection is particularly important to exclude. An opportunistic infection may present identically to an autoimmune-mediated pulmonary disease but the two therapies counteract one another—the former requires a robust immune system and the latter necessitates immunosuppression [7]. Pneumonia inciting a DAH flare requires concurrent treatment [27]. The white blood count may be normal or elevated in both infectious and aPL-related hemorrhage.

Over-the-counter medications, herbal supplements, illicit drugs, toxins, and prescribed medications have been found to induce pulmonary hemorrhage. Culpable medications, (though rarely reported) include propylthiouracil, diphenylhydantoin, amiодarone, mitomycin, D-penicillamine, sirolimus, methotrexate, haloperidol, nitrofurantoin gold, all-trans-retinoic acid, bleomycin, montelukast, zafirlukast, inhaled cocaine, hydralazine, and infliximab [10••, 28]. Thus, it is important to review patients’ medications for offending agents that may accelerate DAH in aPL-positive patients.
During an initial presentation of DAH, a thorough work up to evaluate for other connective tissue disorders and systemic vasculitis should be undertaken. Cytoplasmic and perinuclear anti-neutrophil cytoplasmic antigens (ANCA) (anti-proteinase 3 [PR3] and anti-myeloperoxidase [MPO]) can help evaluate for granulomatosis with polyangiitis, microscopic polyangiitis, and eosinophilic granulomatosis with polyangiitis. Antinuclear antibodies (ANA), double-stranded DNA (dsDNA), complement 3 and 4 levels, urinalysis, creatinine, and anti-glomerular basement membrane antibodies (anti-GBM) can help evaluate for SLE and Goodpasture’s syndrome [29]; however, at times, lupus pneumonitis may be indistinguishable from DAH [30].

Cardiac echocardiogram should be obtained to exclude heart failure given the dyspnea and bilateral ground glass opacities resemble pulmonary edema, and to assess for valvular pathology. Additional imaging to exclude pulmonary embolism is often performed. Pulmonary function tests (PFTs) are rarely performed due to the difficulty patients with dyspnea have in completing the necessary maneuvers. However, if done, PFTs may show restriction; and the diffusing capacity for carbon monoxide (DLCO) may be increased due to blood in the alveolar spaces, which can absorb additional carbon dioxide [10••, 31].

Bronchoscopy with bronchoalveolar lavage (BAL) is critical in patients with APS who present with respiratory failure, dyspnea, or pulmonary infiltrates to diagnose or exclude DAH and infection. Persistent and increasing bloody returns on three sequential BAL sample aliquots support a diagnosis of DAH. Neutrophilic predominance and high numbers (greater than 20%) of hemosiderin-laden macrophages are present on BAL cytology. Of note, increasing bloody returns may precede the hemosiderin-laden macrophages during acute hemorrhage. Yet these macrophages may persist in BAL fluid weeks to months after alveolar erythrocytes have vanished [31]. Lavage specimens should be sent for bacterial, mycobacterial, and fungal stains and cultures, fungal markers, and viral polymerase chain reaction (PCR) tests. An extensive list of infectious pathogens is reviewed elsewhere, but includes Pneumocystis jiroveci, Legionella, Mycoplasma, Nocardia, Leptospirosis, influenza A and B, respiratory syncytial virus, and adenovirus [32]. Transbronchial biopsies during bronchoscopy are not usually recommended [10••]. Surgical lung biopsy (thoracoscopic or open) is not routinely performed given the risk-benefit ratio; however, surgical biopsy may be indicated in certain circumstances when there is persistent diagnostic uncertainty.

Pathologic results of lung biopsy may reveal either pulmonary capillaritis, bland hemorrhage, or diffuse alveolar damage. Pulmonary capillaritis is particularly associated with autoimmune or inflammatory diseases; in a review of 34 lung biopsies of diffuse pulmonary hemorrhage patients (vasculitis- and connective disease-associated DAH in 14 and four cases, respectively), capillaritis was present in 88% of the specimens; however, none of these patients had APS [33]. Causes of bland alveolar hemorrhage include coagulopathies, mitral stenosis and regurgitation, congestive heart failure, and drug-induced; also, it has been reported in APS patients with DAH [10••, 34, 35]. Diffuse alveolar damage is the classic histology observed in ARDS but also evident in infections such as Legionella, Mycoplasma, and Pneumocystis jiroveci, lymphangioleiomyomatosis, radiation therapy, bone marrow transplant complications, and APS-associated DAH [10••].

What Is the Optimal Management of Diffuse Alveolar Hemorrhage in Antiphospholipid Syndrome?

In addition to supportive care, patients are treated with high-dose corticosteroids to induce remission during the acute phase of hemorrhage. Anticoagulation has not been shown to be beneficial in preventing or treating DAH in these patients. Anticoagulants are usually temporarily discontinued during the initial bleeding episode and cautiously restarted depending on the patient’s clinical stability. Many patients require a steroid-sparing agent to achieve remission. Cyclophosphamide, rituximab, mycophenolate mofetil, azathioprine, plasmapheresis, and intravenous immunoglobulin (IVIG) have been used with varying degrees of success (further discussed in the “Literature Review” section). Our management approach is also described in greater detail in the “Our Experience” section.

Literature Review

Methods

We identified 91 APS cases with DAH (primary or associated with another systemic autoimmune disease, with or without catastrophic APS) through a literature review (PubMed Medline search; English only) of publications between January 1991 and December 2018 (eight case series and 15 case reports). PubMed search keywords included: “antiphospholipid,” “alveolar hemorrhage,” “pulmonary hemorrhage,” “catastrophic antiphospholipid syndrome,” “DAH,” and “pulmonary capillaritis.” References of prominent articles were also reviewed.

Catastrophic APS was present in 12 cases; we only included their biopsy results in the assessment of pathology findings but otherwise excluded those patients from the analysis (as this review is not focused on CAPS management) [22, 23, 36–45]. A case series of 13 patients did not report individual treatment and remission data, and thus, these patients were excluded from the outcome analysis; however, we reported their outcomes separately [25].
While assessing the treatment response in the remaining 66 cases, we defined “remission” or “response” as no or less than 10 mg prednisone daily (or equivalent corticosteroid regimen) to maintain remission. For patients who received multiple immunosuppressive medications started non-simultaneously, we included each medication (excluding corticosteroids) in our analysis individually (positive response versus no response). For a small number of patients who received multiple immunosuppressive medications started simultaneously, we included the combination in our analysis (positive response versus no response, excluding corticosteroids). Because we used the most current and final information provided to gauge remission or treatment response, deceased patients were not included as treatment responders. When the data was incomplete, we exercised clinical judgment to best categorize the cases.

Demographics

Seventy-nine aPL-positive patients with DAH (excluding CAPS patients) were mostly males (n = 59; 62%), often Caucasian in their fourth or fifth decade of life (mean age 48 ± 14), suggesting the possibility of a genetic disposition. Eleven patients (14%) had SLE; however, some case series excluded patients with SLE [25, 30, 46]. Eleven of 17 patients in a small case series were current or former smokers [5••]. In a retrospective study of 63 APS patients, compared to APS patients without DAH, patients with DAH were more likely to have mitral valve disease, pulmonary hypertension, skin involvement (livedo reticularis/racemosa or chronic ulcers), obstetric morbidity, and central nervous system disease [25].

Clinical Presentation

Diffuse alveolar hemorrhage was the first presentation of APS in 9/79 (11%) patients [15••, 21, 49–52]; three out of 17 patients in one case series were diagnosed with APS only after they presented with DAH [5••]. Yet, two other reviews of 18 and 13 patients noted a median DAH onset of 5.9 and 5.8 years, respectively, after the diagnosis of APS [25, 46]. Furthermore, many patients have a delay in diagnosis [53, 54], e.g., a 26-year-old female visited her primary care doctor and emergency room for multiple episodes of hemoptysis throughout 1 year before the eventual diagnosis of APS [49].

Triggers

Viral upper respiratory tract infections or bacterial pneumonia were both reported to incite episodes of alveolar hemorrhage [15••, 27]. While periods of supratherapeutic international normalized ratio (INR) could pre-dispose patients to alveolar hemorrhage, some patients had alveolar hemorrhage while off anticoagulation, and the vast majority of published cases revealed sub-therapeutic or therapeutic INR on presentation. Four patients did develop recurrent DAH with a supratherapeutic INR and one patient’s first episode of DAH began 3 weeks after starting warfarin [15••, 25, 27, 46]. Half of patients (52%) were on anticoagulation (typically warfarin) either before or after the diagnosis of DAH, but true prevalence of anticoagulation use is likely higher than reported given the limitations of these studies.

Pathology

Pulmonary capillaritis was seen more frequently than microvascular thrombosis. Half of all published cases with a tissue biopsy (n = 31) demonstrated pulmonary capillaritis (n = 15) [5••, 15••, 24, 25, 38, 41, 46••, 53, 55–57] (Table 1). Micro-thrombosis was present in only seven of 31 biopsies (23%), and five of the seven concomitantly displayed capillaritis [24, 25, 41, 46••, 54, 56, 57]. Interestingly, despite extensive microvascular thrombosis on biopsy, large vessel pulmonary emboli were absent in all seven patients. Bland alveolar hemorrhage or diffuse alveolar damage with or without the presence of mild interstitial inflammation was seen in 45% of biopsies [3, 5••, 25, 30, 36, 45, 46••, 50, 58].

Outcomes (Remission, Recurrence, and Death)

Across the 66 cases included in our outcomes analysis, 43 patients (65%) ultimately achieved remission, 19/43 (44%) with corticosteroids or supportive care alone (Fig. 1), and 24/43 (55%) with immunosuppressive agent(s). At one institution, 10 (59%) of 17 patients achieved complete and sustained remission; interestingly, four of these 10 patients had a mild form of DAH and reached remission without even corticosteroid treatment (details not reported). In the same case series, three (18%) patients relapsed within a median time of 9 months [5••]. Another large case series (n = 18) demonstrated poorer remission rates: 39% achieved complete remission and 28% were alive but remained on high-dose corticosteroids after 1 to 2 years follow up; 72% of all patients experienced recurrent disease [46••].

Based on our review, 36 of 66 patients (55%) experienced recurrent disease, which may underestimate the true relapse rate, since many groups did not report long-term follow up (mean follow up was 35 months; however follow up data was not provided for 11 patients). Symptoms often reappeared during corticosteroid tapering, and many patients were never able to discontinue prednisone.

We found the mortality rate among all cases as 21%. This was similar to published rates in patients with DAH secondary to numerous etiologies: 25% during an acute episode and 16% among those who survived their initial hospitalization [59]. Reported mortality in SLE-associated DAH has ranged widely from 12% in two studies up to 36%, 42%, and 68% in others [60–64]. Data were limited but similar in primary
APS, however, ranged between 23 and 33% [5, 25, 46]. Cause of death was often described as due to “uncontrolled DAH”; however, massive hemorrhage was only rarely described [46]. Documented causes of death included sepsis, respiratory failure due to pulmonary infarction and pneumonia, stroke, spontaneous intracranial hemorrhage, and pulmonary embolism [5, 24, 25, 30, 46, 53, 65].

**Treatment**

Table 2 illustrates major therapeutic interventions utilized and subsequent clinical response. In general, cyclophosphamide- or rituximab-based regimens achieved the highest remission rates (approximately 50%), which aligns with the largest case series reported to date [46]. Intravenous immunoglobulin (IVIG) and plasma exchange together with other immunosuppressive agents demonstrated similar treatment responses (approximately 30%), whereas mycophenolate mofetil and azathioprine use resulted in lower remission rates (approximately 15%).

Given the efficacy of cyclophosphamide in granulomatosis with polyangiitis and other pulmonary vasculitides, this medication was generally utilized initially in the management of aPL-associated DAH [31]. Patients unable to achieve remission with cyclophosphamide ultimately died, remained on high-dose corticosteroids, or responded to rituximab or IVIG [15, 24, 27, 46, 66].

Recent papers have shown success with rituximab, an anti-CD20 monoclonal antibody that targets a surface protein expressed on B-lymphocytes, the precursor of aPL-producing plasma cells. For instance, the first patient with primary APS treated with rituximab was a 32-year-old male who had been hospitalized 15 times over 18 months for recurrent DAH; after unsuccessful treatment with azathioprine, IVIG, cyclophosphamide, and plasmapheresis, two doses of rituximab (1 g each 2 weeks apart) led to a sustained remission over 2 years of follow up [66]. Others have reported similar success with follow up ranging from 1 to 8 years [46, 47].

Several authors have achieved remission using IVIG, both acutely to control an active disease flare and long-term with monthly maintenance infusions. For instance, Janowiak et al.

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**Table 1** Pathology of 31 lung biopsies in antiphospholipid syndrome patients with diffuse alveolar hemorrhage

| Pathology                                           | N (%) |
|-----------------------------------------------------|-------|
| Pulmonary capillaritis                              | 10 (32%) |
| Microvascular thrombosis                           | 2 (6%) |
| Both pulmonary capillaritis and microvascular thrombosis | 5 (16%) |
| Bland alveolar hemorrhage with/without nonspecific/interstitial inflammation, or diffuse alveolar damage (no capillaritis or thrombosis) | 14 (45%) |

*a Only 31 of the 91 reported cases underwent lung biopsy

*b Findings suggestive of small vessel vasculopathy (myointimal thickening/remodeling, thickened arteriole walls, and increased fibroblasts) were mentioned in three biopsies [46, 54, 57]
described a patient with six recurrent DAH episodes and multiple infections who only remained disease-free after the addition of monthly IVIG to mycophenolate and prednisone [27]. Others report similar success with IVIG [15, 51]; however, some patients did not achieve remission with IVIG [46, 65, 66]. Plasma exchange, which is traditionally used in CAPS, has been effective in a third of patients with DAH without CAPS when used in combination with other immunosuppressive agents [5, 50, 57, 67, 68].

Monotherapy with mycophenolate mofetil had a relatively low response rate [15, 27, 46, 47, 68]; however, the combination of rituximab and mycophenolate mofetil resulted in complete remission in two patients [46]. None of the six patients on monotherapy azathioprine in the largest case series attained remission [46], and others reported similar outcomes [25, 30, 50, 53].

Hydroxychloroquine was used as monotherapy in only two cases (one resulted in remission and the other treatment failure) and demonstrated intermittent success when utilized as an adjunct agent [5, 46, 47, 66, 68].

Our Experience

Diagnostic Considerations

In patients with clinically significant aPL profiles (persistently positive LA test and/or moderate-to-high titer aCL/aβ2GPI IgG/M), when imaging studies are suggestive of DAH, bronchoscopy is important during initial or ambiguous presentations. During the initial diagnostic assessment, we also obtain chest CT with contrast as pulmonary embolism can present with hemoptysis. In patients with APS and recurrent, well-documented DAH and without suspicion for additional underlying disease or infection, we generally do not perform repeat bronchoscopy.

We do not routinely recommend lung biopsy due to associated risks in patients who are generally anticoagulated and at high risk for thrombosis. Lung biopsy is considered if the etiology of the DAH is unclear from the serologic work up and an accurate diagnosis cannot otherwise be made. If performed, the pathologist should perform immunofluorescent staining to assess for immune-complex and complement deposition, and tissue cultures should be sent as reviewed above. Of note, if the clinical scenario and laboratory tests suggest APS, the absence of capillaritis or microthrombosis on biopsy should not preclude the APS diagnosis.

Minor hemoptysis presents a diagnostic dilemma when it develops in a patient with APS on warfarin. Is the warfarin to blame? We urge providers to consider DAH on the differential to avoid missed and delayed diagnoses. And in those with documented APS, high clinical suspicion for DAH is necessary, as documented cases of hemoptysis incorrectly attributed to warfarin overdose rather than a first occurrence of alveolar hemorrhage have been reported [27]. In addition, when patients do not yet carry the DAH diagnosis, symptoms may be falsely attributed to respiratory infections.
**Therapeutic Considerations**

Given the rarity of the disease, lack of controlled studies, limited literature regarding treatment, and heterogeneity, there is no “one size fits all” approach to treatment. In our experience, two different presentations of aPL-positive patients with DAH exist: (a) acute—with moderate to severe pulmonary hemorrhage (with varying degrees of respiratory failure) requiring hospitalization and (b) chronic—with mild hemoptysis, dyspnea, and/or positive imaging findings, usually seen as an outpatient. Patients in both settings often require steroids for disease control and a steroid-sparing immunosuppressive for maintenance therapy, as we discuss below.

In hospitalized patients, supportive care is a mainstay of acute management, ranging from supplemental oxygen to mechanical ventilation, along with as-needed transfusions of blood products. Empiric antibiotic therapy is often co-administered with corticosteroids at initial presentation and subsequently if a patient clinically decompensates. Of note, immunosuppressed APS patients with DAH are especially susceptible to pulmonary infections (typically bacterial), which can manifest with hemoptysis. We do not routinely treat with antiviral or antifungal antibiotics unless laboratory or culture data suggest a particular pathogen. High-dose corticosteroids (intravenous methylprednisolone 500–1000 mg daily for 3 days, followed by oral prednisone 1 mg/kg daily in divided doses (maximum 60 mg day)) are generally the primary treatment during the acute phase. Due to high mortality in these patients, corticosteroids are rapidly initiated even when diagnostic tests are still pending. For patients with life threatening or ventilator-dependent DAH, usually in the setting of CAPS, we also use plasma exchange and/or intravenous immunoglobulin [69].

Anticoagulation therapy is temporarily discontinued during the acute severe bleeding episode; however, the risks and benefits of stopping anticoagulation should be carefully calculated in aPL-positive patients with/without history of thrombosis. When stopped, we recommend restarting anticoagulation as soon as the acute bleeding is controlled. In cases with concurrent pulmonary embolism and hemorrhage, starting anticoagulation and corticosteroids simultaneously may be required [70]. A multidisciplinary team approach should appraise the risks and benefits, i.e., timing and dosage of safe anticoagulation [71]. In aPL-positive patients without a history of thrombosis presenting with DAH, anticoagulation does not treat or prevent DAH and we do not recommend anticoagulation for treatment purposes. However, we recommend continuing prophylactic dose anticoagulation to prevent thrombosis in hospitalized patients.

In an outpatient setting with mild DAH in the absence of respiratory compromise, we recommend initiating oral prednisone (or increasing the dose to) 1 mg/kg daily to a maximum of 60 mg daily. As discussed above, in patients with suspected new-onset DAH, urgent referral to a pulmonary team for confirmatory bronchoscopy is required. For patients with known DAH and recurrent albeit mild symptoms, oral corticosteroids with close follow up is often sufficient.

Both in an inpatient and outpatient setting, in addition to steroids, we recommend starting steroid-sparing immunosuppressive agents as, based on our experience, the risk of recurrence is relatively high during corticosteroid tapering. Our first-line therapy is generally mycophenolate mofetil (quickly accelerating the dose to 3 g/day) with or without rituximab (1000 mg twice 2 weeks apart), once infectious etiologies have been excluded. We add rituximab to mycophenolate mofetl especially in aPL-positive patients with hematologic and/or other microthrombotic manifestations, e.g., thrombocytopenia or livedoid vasculopathy, and also in patients hospitalized with relatively severe hemorrhage. After the DAH is controlled, we decrease the prednisone dose with close monitoring, usually by 10–15 mg per month. We do not use cyclophosphamide as first-line therapy due to the high incidence of treatment-associated toxicity.

In cases of recurrent DAH on mycophenolate mofetil, we add rituximab depending on the other aPL-manifestations discussed above. No data exist on the long-term management of rituximab in APS patients with DAH; we generally do not continue rituximab if there is full recovery. If patients have no response or recurrence of DAH despite the combination of mycophenolate mofetil and rituximab, we next add IVIG. We have treated several patients who achieved remission with the addition of IVIG 2 g/kg over 4–5 days, every 1–3 months, to mycophenolate mofetil (with or without a history of rituximab use) (unpublished data).

In all aPL-positive patients with DAH, despite the lack of clinical data, we add hydroxychloroquine (200–400 mg daily) and a statin (e.g., atorvastatin 10-20 mg/day) to the treatment regimen. Hydroxychloroquine reduces the risk of thrombosis in experimental aPL/APS models and lupus patients; similarly, statins ameliorate the proinflammatory and prothrombotic markers in aPL-positive patients [72–75].

**Prognostic Considerations**

We do not understand why certain patients experience a single episode that clears with corticosteroids, while others require long-term immunosuppressive agents. Based on our experience, patients may continue to have abnormalities on imaging despite resolution of symptoms. Some of these patients, whose symptoms of DAH have resolved, but have persistent opacities on imaging, may have recurrence of mild hemoptysis during upper respiratory tract infections or when their INR is supratherapeutic. In asymptomatic patients with persistent imaging findings, we continue the immunosuppressive agents while tapering and ultimately discontinuing prednisone as discussed above.
Monitoring for recurrence can be challenging, particularly when patients are asymptomatic with normal laboratory findings. For patients on chronic immunosuppression for prior hemorrhagic episodes, we recommend obtaining chest radiograph and pulmonary function tests including DLCO yearly or with changes in respiratory symptoms.

In our experience, patients with DAH (without CAPS) have lower mortality rates than those reported in the literature; we attribute this to publication bias for this rare condition.

Conclusion

Diffuse alveolar hemorrhage is a rare yet potentially fatal complication of APS, in which red blood cells leak from the alveolar capillaries into the intra-alveolar space. Pulmonary capillaritis, along with antibody-mediated endothelial proliferation, likely contributes to its pathogenesis, yet is not always observed pathologically. Diffuse alveolar hemorrhage typically presents with hypoxemia and subsequent respiratory failure, anemia, hemoptysis, and diffuse radiographic pulmonary infiltrates.

Given the rarity of pulmonary hemorrhage in APS, no randomized trials exist. Treatment is guided by case reports, case series, and inter-disciplinary experience. While anticoagulation is standard of care for the treatment and prevention of thrombotic complications of APS, its use is contraindicated during acute DAH episodes and is not efficacious in preventing DAH. In addition to supportive care, high-dose corticosteroids are the initial treatment for DAH in APS. The addition of thymoglobulin or rituximab as a first-line treatment strategy, and adding IVIG, plasmapheresis, and cyclophosphamide in patients with limited response.

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