How Not to Be Misled by Disorders Mimicking Angioedema: A Review of Pseudoangioedema

Michelle Fog Andersen a, Hilary J. Longhurst d, Eva Rye Rasmussen b, Anette Bygum c

a Department of Otorhinolaryngology, Head and Neck Surgery, University Hospital of Sjaelland, Koege, b Department of Otorhinolaryngology, Head and Neck Surgery, University Hospital of Copenhagen, Copenhagen, and c HAE Centre Denmark, Department of Dermatology and Allergy Centre, Odense University Hospital, Odense, Denmark; d Department of Immunology, Barts Health NHS Trust, London, UK

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Abstract
Background: Angioedema is a vascular reaction involving the lower dermis, subcutis and/or submucosal tissue and causing a temporary localized swelling in any part of the body. For many health care professionals, the diagnosis presents an ongoing challenge; several disorders may manifest with subcutaneous or submucosal swelling and falsely be assumed to be angioedema. The clinicians at the emergency department and in the immunology/allergy clinics must be skilled at recognizing the features of angioedema and its differential diagnosis. Methods: The review is based on a literature search with specific indexing terms in PubMed, a review of bibliographies and the authors’ clinical experience. Results: The most essential diseases that mimic angioedema, the so-called pseudoangioedemas, will each be discussed and illustrated by clinical photos, pointing out key features that help clarify the diagnoses and differentiate these from classic angioedema. Conclusions: A variety of dermatologic conditions can cause swelling that resembles angioedema, some with a potentially fatal outcome if misdiagnosed. Knowledge of pseudoangioedema is fundamental in the emergency setting when handling patients with edema and should be kept in mind when assessing an atypical angioedema case.
Angioedema is a clinical symptom defined by local swelling of the deeper layers of the skin, mucous membranes, or both, including the gastrointestinal tract and upper airway. Although the swelling can affect any part of the body, it has a predilection for areas with loose connective tissue, including the face (especially the lips, tongue and periorbital area), pharynx, larynx, abdomen, genitalia and extremities [11–13]. The swellings manifest as recurrent episodes of pronounced localized edema with ill-defined margins. Unlike other forms of edema, angioedema is nonpitting, often asymmetric and has a tendency not to involve gravitationally dependent areas. The skin is usually normal in color, but can be slightly erythematous. The majority of patients with angioedema describe a tingling, slightly numb or even burning sensation of the affected site, while itching is not a typical symptom. The swelling is typically of slow onset over several hours and lasts for <72 h, resolving spontaneously without staining or desquamation of the skin. However, angioedema can have a very rapid onset when it occurs due to anaphylaxis. Over half of patients have concomitant urticaria [11, 12].

Pseudoangioedema

There are several conditions presenting with subcutaneous swelling that superficially resemble angioedema. Based on key signs and symptoms (online suppl. table 1; see www.karger.com/doi/10.1159/000445835), these so-called pseudoangioedemas can clinically be distinguished from true angioedema. The most common and important diseases that can mimic angioedema are listed below.
Acute Contact Dermatitis

Acute contact dermatitis is an inflammatory response of the skin induced by contact with a foreign substance. The reaction can be triggered by direct interaction with an irritating agent (irritant contact dermatitis) or allergenic chemical substance (allergic contact dermatitis) causing a type IV allergic reaction [14]. Acute contact dermatitis of the face is often misdiagnosed as angioedema, as it can cause severe swelling of the facial and periorbital skin, especially after contact with hair dye (fig. 1) [14, 15]. The reaction can become so severe that intubation may be necessary. It can be distinguished from angioedema and other pseudoangioedemas by demonstration of superficial erythema, dermatitis, prominent pain or pruritus and a history of exposure to a foreign substance. Finally, the skin will often peel as the swelling resolves, in contrast to patients with angioedema [14]. Antihistamines are ineffective, but symptoms respond to corticosteroids given topically or systemically. Diagnosis is verified by patch testing and the management involves strict avoidance of any contact allergens identified, along with withdrawal of the corticosteroids once symptoms are well controlled.

Drug Rash with Eosinophilia and Systemic Symptoms

Drug rash with eosinophilia and systemic symptoms (DRESS) refers to an uncommon but rather severe adverse drug-induced skin reaction. Among a wide list of drugs associated with this reaction, aromatic anticonvulsants (phenytoin, phenobarbital and carbamazepine) are the most common [16]. DRESS may mimic angioedema due to its clinical manifestation with facial or more widespread rash (fig. 2) [17]. Other characteristic findings include fever, eosinophilia, lymphadenopathy and internal organ involvement, mainly the liver and kidneys, which differentiate this condition from angioedema. The onset of the disease is usually seen within 6 weeks after the initiation of drug exposure and DRESS has a longer duration than most allergic drug reactions [16, 18]. In fact, the symptoms can even exacerbate despite discontinuation of the drug. A history of the patient’s medication, together with objective findings and blood tests, is therefore of utmost importance. Traditional patch testing cannot be used to confirm the diagnosis, which has to be clinical. Management consists of discontinuation of the causative drug. Most patients recover completely after drug withdrawal, but treatment regimens with corticosteroids are commonly used as well [18].

Dermatomyositis

Dermatomyositis is a common idiopathic inflammatory myopathy that affects both skeletal muscle and the skin [19]. Although the etiology is poorly understood, an immunological pathogenesis has been suggested. The disease is characterized by proximal muscle weakness and cutaneous eruption in the form of erythema. The most common cutaneous features include the heliotrope rash: a distinctive reddish-purple erythematous rash around
the eyes and Gottron’s sign or papules [19, 20]. It may present with periorbital edema, which can resemble that of angioedema (fig. 3). However, the presence of symmetrical proximal myositis, the cardinal muscular feature, along with the characteristic cutaneous lesions, fatigue, weight loss and fever, distinguish this disease from angioedema [11]. Clinical suspicion is confirmed by biochemical, electromyographic and histological evidence of inflammation, with raised serum creatine kinase, antisynthetase antibodies, myopathic features on electromyography and magnetic resonance imaging and typical muscle or skin biopsy features which may be patchy. Management, usually within a specialist center, involves immunosuppression [21].

**Morbus Morbihan**

An uncommon cutaneous disorder called Morbus Morbihan, also known as persistent edema of rosacea, is a skin condition which can be confused with angioedema [22]. It is considered a rare complication of rosacea and is characterized by persistent erythematous edema restricted to the forehead, glabella, upper eyelids and cheeks (fig. 4). The edema worsens gradually over months to years with a solid consistency. The patient has no other symptomatic complaints and neither specific laboratory nor histopathologic findings have been observed [23]. The chronic nature of the skin condition, together with the restricted location, should differentiate this from angioedema. There is no guideline for its management. However, reported therapy includes long-term systemic corticosteroids and/or oral antibiotics such as doxycycline and isotretinoin [24].

**Superior Vena Cava Syndrome**

Superior vena cava syndrome is a group of symptoms caused by obstruction and thereby impaired blood flow through the superior vena cava into the right atrium. In most cases, the obstruction is caused by a malignant tumor within the thorax. The clinical manifestations of superior vena cava syndrome usually develop slowly and include dyspnea, cough and hoarseness. Because the drainage of blood to the heart is obstructed, the syndrome can, in the early stages, masquerade as angioedema, due to a gradual development of edema in the face and upper extremities (fig. 5) [25, 26]. Clues to differentiate the 2 diagnoses include vein distension across the chest and neck as well as an increase of signs when the patient is in a supine position [11, 27]. X-ray or CT scan of the chest including the thoracic inlet usually confirms the diagnosis and further management is then undertaken by the thoracic oncology team.

**Hypothyroidism**

Insufficient production of thyroid hormones by the thyroid gland causes hypothyroidism, which is most often autoimmune [28]. The condition presents with a wide array of symptoms including weight gain, constipation, dry skin, thinning of hair, hoarse voice, fatigue, lethargy,
Disorders Mimicking Angioedema: Pseudoangioedema

Pseudoangioedema

Depression and cold intolerance. Severe hypothyroidism can manifest with puffiness of the face and lips very similar to angioedema (fig. 6). When generalized nonpitting edema (myxedema) becomes a manifestation, periorbital edema is often seen as a symptom. Nonetheless, it is not transient like angioedema [10, 17]. Identification of the diagnosis is based on clinical features and low levels of thyroid hormones. The treatment of choice for hypothyroidism is thyroxine replacement [28].

Subcutaneous Emphysema

Air bubbles or other gases trapped in the subcutaneous tissues, namely subcutaneous emphysema, cause a sudden onset of swelling in the affected area [29]. Air can migrate through the various fascial planes to involve the thorax, abdominal wall, perineal region, extremities and most often the neck or face (fig. 7), causing the condition to mimic angioedema. The air may become trapped as a result of surgery or trauma, or can occasionally develop spontaneously. Subcutaneous emphysema can usually be diagnosed clinically by crepitus, a characteristic crackling sensation created as the gas is pushed through the tissue during palpation [29, 30]. In patients with this significant finding, the differential diagnosis of angioedema is ruled out. In doubtful cases, an X-ray or CT scan could be performed to illustrate the air beneath the skin surface and to identify the source of the emphysema.

Orofacial Granulomatosis

Orofacial granulomatosis represents a group of chronic diseases affecting the soft tissues of the oral and maxillofacial region secondary to an underlying granulomatous inflammation [31]. The clinical presentation commonly shows persistent swelling of the lips (fig. 8). The group includes Melkersson-Rosenthal syndrome, an idiopathic disorder represented by a classic triad of persisting lip or facial swelling, facial nerve paralysis and fissured dorsal tongue (lingua plicata). Monosymptomatic cases with labial involvement alone are referred to as cheilitis granulomatosa. The orofacial edema is painless and asymmetrical, most often affecting the upper lip. Clinically, it is similar to angioedema and plays an essential role as a differential diagnosis [10, 31, 32]. However, its chronic nature should distinguish it from typical angioedema. The etiology remains unclear but has been linked to an abnormal immune reaction. Diagnosis is confirmed by histology. Management often requires the use of corticosteroids or immunosuppression [32].
Hypocomplementemic Urticular Vasculitis Syndrome

Hypocomplementemic urticarial vasculitis syndrome is a form of severe, cutaneous, small-vessel vasculitis characterized by urticaria and abnormally low levels of complements C1q, C3 and C4 with C1q antibodies [33, 34]. The most distinctive manifestation of this rare disease is recurrent episodes of chronic, nonpruritic, urticarial skin lesions (fig. 9) associated with systemic involvement including pulmonary disease, abdominal pain, leukocytoclastic vasculitis, arthritis, arthralgia and glomerulonephritis. Underlying malignancy, infection or connective-tissue disease should be excluded. In addition, angioedema is the initial clinical presentation in over half of patients, often involving the facial area and upper extremities – a clinical picture found to misguide less-experienced physicians [33, 35]. However, the characteristic lesions in hypocomplementemic urticarial vasculitis syndrome are typically painful and often resolve with postinflammatory hyperpigmentation or purpura upon their resolution. These characteristics, including the extracutaneous and systemic involvement, are all uncommon for typical angioedema. The appropriate management is determined by the severity of the disease, and may include combinations of antihistamines, hydroxychloroquine, corticosteroids and immunosuppression [36].

Clarkson’s Disease

Systemic capillary leak syndrome, also called Clarkson’s disease, is a life-threatening condition characterized by recurrent episodes of sudden hypovolemic shock and massive edema due to the capillary leakage of plasma from the intravascular to the extravascular compartments [37, 38]. The pathogenesis is still unclear, but immune dysregulation may play an essential role. Systemic capillary leak syndrome is diagnosed clinically after exclusion of other diseases caused by systemic capillary leak. A diagnosis of angioedema should be considered upon the initial presentation. Nonetheless, unlike angioedema, the cutaneous swelling of systemic capillary leak syndrome is generalized and symmetrical (fig. 10). Moreover, the rapid shift results also in hypovolemia, hemoconcentration and reduced serum albumin – a triad not characteristic of angioedema. Acute treatment is supportive. Corticoste-
Disorders Mimicking Angioedema: Pseudoangioedema

Gleich’s Syndrome
The syndrome of episodic angioedema with eosinophilia, also known as Gleich’s syndrome, is a rare disorder of unclear etiology characterized by recurrent episodes of idiopathic angioedema, eosinophilia and elevation of the serum immunoglobulin, lasting up to a few months [39]. The clinical picture also demonstrates weight gain caused by fluid retention, fever, pruritus, and, in some cases, urticaria. It is characteristic for the syndrome that there is no systemic organ involvement. The presence of specific laboratory features, together with the other characteristic clinical manifestations, should differentiate this entity from classical angioedema. Although debate surrounds the pathophysiology, increased serum levels of interleukins 5 and 6 has been described [39, 40]. Systemic corticosteroids are used when treatment is required.

Cluster Headache
Cluster headache, also known as histamine headache, is a rare cause of unilateral head or facial pain and periorbital edema, often associated with autonomic features such as conjunctival injection, ptosis, pupil constriction, watering of the eyes or rhinitis (fig. 11). Untreated, symptoms last up to a few hours and may recur. Pain is typically intense and resistant to antihistamines and topical steroids. Oral steroids provide relief. Treatment of episodes is usually with rapid-acting tryptan or high-flow oxygen. Verapamil is the drug of choice for prophylaxis [41]. The presence of the characteristic headache, together with the other clinical manifestations, should differentiate this entity from classical angioedema.

Idiopathic Edema
Idiopathic edema is a self-limited condition of persistent fluid retention, primarily in women [3]. The fluid retention is typically most prominent in premenstrual periods, which is why the condition is also known as ‘cyclical edema’. It is most prominent on the extremities or abdomen after a prolonged upright position and also in the facial area, including pronounced periorbital edema, after recumbency overnight (fig. 12). This distinctive alteration over time, combined with an excessive weight gain from morning to evening and pitting edema, is very uncharacteristic for angioedema. Also, the edema is pitting, in contrast to the nonpitting nature of angioedema. The diagnosis is one of exclusion and should only be considered when there is no evidence of cardiac, hepatic, renal or thyroid disease, all well-known causes of edema. Patients with idiopathic edema often become dependent on diuretics when trying to minimize the fluid retention. It is therefore important to avoid diuretics and, instead, consider angiotensin-converting enzyme inhibitors when treating these patients [42].

Discussion
A variety of dermatologic conditions can cause swelling that resembles the diagnosis of angioedema. This review has discussed the most essential diseases that mimic angioedema, the so-called pseudoangioedemas, and also pointing out key features that help clarify the diagnoses and differentiate these from classic angioedema. Working knowledge of this kind is fundamental in the emergency setting when dealing with patients with edema, and the list of diseases should be kept in mind when assessing an atypical angioedema case. In this way, determination of the root cause of symptoms can be made, leading to more effective and appropriate management of sometimes life-threating conditions.
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References


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