Paracetamol and Chlorpheniramine Combination Linear IG A Bullous Dermatosis

Parasetamol ve Klorfeniramin Kombinasyonuna Bağlı Lineer IG A Büllöz Dermatozu

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ABSTRACT

Linear IG A Bullous Dermatosis (LABD) is a rare autoimmune, subepidermal vesiculobullous dermatosis. The etiology is unknown in most patients. Most recent case reports report drug-induced LABD cases. 35 year - old male patient applied to our clinic with the complaint of tense bullous lesions located on trunk and extremities which started to occur after using a combined drug including paracetamol and chlorpheniramine. The lesions on the patient had responded quickly to systemic and topical corticosteroid treatment, no new lesions had existed since fifth day of treatment and older lesions had begun to be epithelized. Clinical findings and histopathologic and direct immunofluorescent examination findings are important for differentiation between LABD and other bullous dermatoses. The clinical course is lighter than the idiopathic form in drug-induced LABD. Drug-induced LABD is rarely observed compared to idiopathic form. Many drugs can cause this condition. The reported drug lists and variety continue to increase. Dermatologists should be aware of the rare adverse effects that may occur as a result of drug use.

Keywords: Linear IG A bullous dermatosis; paracetamol; chlorpheniramine

ÖZET


Anahtar Kelimeler: Lineer IG A büllöz dermatozu; parasetamol; klorfeniramin.
lieve that our case was the first LABD case report induced by paracetamol + chlorpheniramine.

CASE REPORT

A 35-year-old male patient was admitted to our outpatient clinic with the complaint of vesiculobullous lesions on the trunk and extremities about 1 week ago. He was accompanied by itching in together. It was learned that the rashes of the patient had been intermittent for 2 years and regressed with topical steroid treatment. There was no significant feature his history and in family history. He had no history of drug use except for a combination of paracetamol + chlorpheniramine which he used intermittently for headache. He had recently used a drug with a combination of paracetamol + chlorpheniramine 6 days before the onset of his complaints. During the dermatological examination, diffuse, scattered, sharp-edged, on annular erythematous plaques, strained bullous lesions and diffuse erode areas were observed in the trunk and extremities of the patient (Figure 1-4). Nikolsky’s sign was negative. Examination of the oral mucosa and other mucosal areas was normal. The patient’s vital signs were stable. There was no significant findings except for leukocytosis (WBC = 21500/µl), sedimentation (45 mm/h) and CRP elevation (CRP=33.4 mg/dl) in the laboratory findings. Biopsy was taken for histopathological and direct immunofluorescence examinations with the preliminary diagnoses of bullous pemphigoid, LABD, dermatitis herpetiformis, epidermolysis bullosa acquisita and generalized fix drug eruption. During Histopathological examination; In the epidermis, It was observed

![Figure 1-4: During the dermatological examination, diffuse, scattered, sharp-edged, on annular erythematous plaques, strained bullous lesions and diffuse erode areas were observed in the trunk and extremities of the patient.](attachment:image)
septant orthokeratosis acanthosis, spongiosis, large bulla formation which is separating the dermo-epidermal juncture in the subepidermal area and containing mixed types of inflammatory cells. Inflammatory cells were eosinophil-containing polymorphonuclear leukocytes and lymphocytes and were also found in the perivascular and interstitial areas. Direct immunofluorescence examination of the perilesional skin revealed linear IgA, IgG and C3c accumulation in the basement membrane in accordance with LABD. After biopsy, the patient was treated with 40 mg/day systemic steroid, topical steroid creams and topical antibiotic creams for erode infected lesions. On the 5th day of systemic steroid treatment, no new lesion emerged, and the patient’s existing lesions were completely epithelized by responding dramatically to the treatment (Figure 6-9).

DISCUSSION

LABD is an autoimmune, subepidermal vesiculobullous dermatosis. It may occur due to idiopathic or drug. The clinical status is heterogeneous in both. The etiology is unknown in most patients. Most recent case reports report drug-induced LABD cases, however it may be equally common in idiopathic form. LABD-related drugs, most commonly vancomycin, are various antibiotics, nonsteroidal anti-inflammatory drugs (eg, diclofenac, naproxen, piroxicam), lithium, captopril, amiodarone, phenytoin, cyclosporine, furosemide, interferon alpha and somatostatin. Genetic factors may also contribute to the development of LABD. The relationship of LABD with HLA B8, HLA Cw7, HLA DR3, HLA DQ2 and tumor necrosis factor-2 allele has been reported.7

The pathophysiology underlying the drug-induced LABD is still unclear. However, it is thought to result from an immune response against a drug-induced hapten-protein antigen. An alternative explanation for the pathogenesis of drug-induced LABD is the structural modification of protein molecules or the emergence of a previously concealed antigenic determinant that leads to an immune response of the drug.8 Target antigens are different antigenic structures in lamina lucida, sublamina densa or located both regions. Identification of target antigen in LABD is important for clinical picture and prognosis. For example; if the target antigen is Type VII collagen, patients are less probability to respond to treatment and are considered to be a subset of epidermolysis bullosa acquisita. Similarly, patients with antibodies directed against bullous pemphigoid antigens have clinically bullous pemphigoid. However, they indicate an IgA response instead of an IgG response. Those with antibodies to laminin 332 may potentially have mucosal involvement.9,10 In addition, the subunits of laminin 332 have been the target antigen in vancomycin-induced linear IgA dermatosis.11

The clinical presentation of drug-induced cases is generally not different from that of idiopathic patients. They usually show a clinical appearance similar to other vesiculobullous dermatoses such as bullous pemphigoid and dermatitis herpetiformis. Rarely, the lesions may be localized or show similar clinical features such as morbilliform drug eruptions, erythema multiforme or toxic epidermal necrolysis (TEN). The presence of mucosal involvement is variable there is no mucosal involvement in our case.12 The lesions were in the form of tense vesiculobullous lesions with asymmetric distribution in the trunk and extremities.

Most of the drug-induced LABD cases are observed in adults. The disease usually occurs within a
month of the onset of the drug and cures within a few weeks of discontinuation. However, repeated exposure to the causative drug may cause the disease to develop more rapidly. For example; Rash in vancomycin-induced LABD cases; after the first dose, it occurred between 1 and 15 days; it occurred 2 days after the first dose in one case reported to be related to acetaminophen. In our case, lesions appeared 6 days after ingestion of the drug, and all lesions were epithelized by treatment 5 days later after cessation of the trigger drug.

The differential diagnosis of LABD includes dermatitis herpetiformis, bullous impetigo, bullous pemphigoid and acquired epidermolysis bullosa. Direct immunofluorescent examination is the gold standard method to differentiate LABD from other bullous disorders. Especially, bullous pemphigoid which is more common in elderly patients, is differentiated from LABD with the histopathological finding of more prevalent eosinophil infiltration compared with LABD. Also, direct immunofluorescent examination of bullous pemphigoid is characterized with linear IgG and C3 deposition on basement membrane.

Diagnosis is made by histopathological examination and direct immunofluorescence examination which is the gold standard in diagnosis. The histopathological findings of LABD are non-specific and often resemble dermatitis herpetiformis. An infiltration which subepidermal bullae and PMNLs are predominant is characteristic. During direct immunofluorescence examination, linear IgA deposition is typically seen in BMZ. Linear C3 deposition may also be observed. In some patients, both linear IgA accumulation and immunoglobulin G (IgG) accumulation may be seen in BMZ. Immunoglobulin M (IgM) accumulation has rarely been reported. In our case, linear IgA, Ig G and C3c accumulation was seen in the basement membrane zone. Some researchers have suggested that coexistence of linear IgA and IgG accumulations in LABD is coincidental or these cases be categorized as vesicular pemphigoid or bullous...
pemphigoid/LABD overlap or linear IgA / IgG bullous dermatosis.\textsuperscript{15-17} The induction of autoantibody synthesis in autoimmune bullous dermatoses is not well understood. Also, pathogenic anti-IgA, anti-IgG or the factors which are leading to the synthesis of both antibodies are unknown. Recent studies make think that specific cytokine microenvironments (eg transforming growth factor \( \beta_1 \), interleukin 4, interferon gamma) play a role.\textsuperscript{18} Data on treatment options are limited in LABD. The treatment approach is usually based on case reports and case series. The first step in drug-related cases is discontinuation of causative drug and dapsone, sulfapyridine, topical and systemic steroids can be selected for treatment.\textsuperscript{4} The clinical course is milder than the idiopathic form in drug-induced LABD. In drug-induced LABD, cessation of new lesion formation typically occurs within three days after discontinuation of the drug, and complete recovery usually occurs within a few weeks.\textsuperscript{19} In our patient, near-complete epithelialization developed in the lesions 5 days after discontinuation of the causative drug and with systemic steroid treatment.

As a result; Drug-induced LABD is rare compared to idiopathic form. Many drugs can lead to this Picture.\textsuperscript{20-22} The reported drug lists and variety are increasing. Dermatologists should be aware of this rare adverse effect that may occur as a result of drug use.

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**Conflict of Interest**

No conflicts of interest between the authors and / or family members of the scientific and medical committee members or members of the potential conflicts of interest, counseling, expertise, working conditions, share holding and similar situations in any firm.

**Authorship Contributions**

All authors contributed equally while this study preparing.

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**REFERENCES**


