There is good evidence for the comorbidity of psoriasis and bullous pemphigoid (BP) since its first description in 1929 (1, 2). The onset of BP on top of psoriasis usually occurs after the administration of anthralin, coal tar, salicylic acid or phototherapy (3). Treatment of BP in psoriasis frequently poses a therapeutic dilemma, because the treatment of choice for BP is systemic corticosteroid, which is relatively contraindicated in psoriasis. We report here a patient with BP and psoriasis who achieved long-term remission of BP after 2 injections of rituximab, an anti-CD20 monoclonal antibody.

CASE REPORT
An 83-year-old man presented with chronic plaque-type psoriasis, which had been well-controlled with topical treatment and ultraviolet B (UVB) phototherapy for over 10 years. He weighed 105 kg and could ambulate with assistance. In January 2005, some itching vesicles and urticarial plaques developed on his trunk and lower limbs over a period of 2 months. Skin biopsy revealed papillary dermal oedema with some perivascular infiltration of eosinophils. In addition, direct immunofluorescence revealed linear IgG and C3 deposition along the basement membrane (Fig. 1). Indirect immunofluorescence detected anti-basement membrane antibody at a titre of 1:20. A diagnosis of BP was then established. Initially, topical clobetasol propionate ointment and oral prednisolone (40 mg/day) were administered. The dosage of prednisolone was gradually tapered and eventually discontinued one year later. No new lesions were noted in the following 2 years. Methotrexate, 5 mg/week, was added from 2007 as a maintenance dose for psoriasis. In August 2008, the patient had another flare-up of BP, with numerous intense pruritic vesiculobullous eruptions on the extremities (Fig. 2). Due to the pre-existing skin atrophy and osteoporosis from his previous prednisolone use, he received 2 doses of rituximab 500 mg therapy weekly, together with a short course of oral prednisolone (20 mg daily for 2 weeks, then 30 mg every other day for another 2 weeks). Concomitant methotrexate 7.5 mg weekly was given in the first 3 months and the dosage was then tapered to 5 mg weekly. No further vesicles occurred after 6 weeks, and the patient has remained free of BP for more than one year to date. No adverse events were experienced during rituximab treatment.

DISCUSSION
Chronic disruption of the basement membrane layer in psoriasis may result in altered antigenicity and the subsequent production of autoantibodies (4). BP with psoriasis is typically associated with autoantibodies against type XVII collagen (Col17) (5). In addition, a novel 200-kDa protein (p-200) has been detected as another antigen for a subset of autoimmune subepidermal blistering diseases, which is often associated with psoriasis (6).

Systemic corticosteroid is usually the first-line treatment for BP. However, it is also known to trigger the development of erythrodermic or pustular psoriasis once discontinued. Rituximab is an anti-CD20 monoclonal antibody indicated for the treatment of rheumatoid arthritis and non-Hodgkin lymphoma. It is also increasingly used in the treatment of autoimmune bullous disorders, including BP and pemphigus (7). Thus, the use of rituximab in the treatment of BP in psoriasis is an attractive alternative. There are many papers on successful use of rituximab in patients with either psoriasis or psoriatic arthritis (8, 9). However, there is only one case report of the use of rituximab in the treatment of losartan-induced BP in a patient with psoriasis (10). In addition, there are reports of induced or aggravated psoriasis following the use of rituximab for other diseases.

Fig. 1. (a) Linear IgG, and (b) C3 deposition along the basement membrane (a: ×100 and b: ×40).
therapy well without increased risk of infections, corticosteroid-sparing agents, and concomitant systemic corticosteroid is still needed, which might cause further skin atrophy and psoriasis aggravation following its discontinuation after prolonged use (13). Rituximab is mostly used on refractory BP, in particular mucous membrane pemphigoid (14). Although higher infectious complication may exist, compared with azathioprine and mycophenolate mofetil monotherapy most patients tolerated rituximab and psoriasis aggravation following its discontinuation after prolonged use (13). Rituximab is mostly used on refractory BP, in particular mucous membrane pemphigoid (14). Although higher infectious complication may exist, compared with azathioprine and mycophenolate mofetil monotherapy most patients tolerated rituximab therapy well without increased risk of infections, cardiovascular events, malignancies or fatality over time (15). In addition, it has the added benefit of sustained clinical remission and reduced need for systemic prednisolone. Although methotrexate has also been used for the treatment of BP, in our patient, methotrexate had been added for psoriasis control before the flare-up of BP. Thus, we believe that the clinical remission of BP was achieved mainly by rituximab. The use of methotrexate may have had only a minor additive role. In addition, oral prednisolone was tapered and discontinued in only one month, a much shorter time than the one-year course of oral prednisolone treatment in 2005. Although the possible long-term side-effects and the high costs of rituximab compared with conventional treatment do not support its generalized use for BP, rituximab may offer a valuable therapeutic alternative for selected cases of BP, such as BP in psoriasis. Further large-scale study is needed to evaluate the efficacy and safety of rituximab in treating this group of patients.

REFERENCES

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Fig. 2. Clear-fluid-containing, 1–1.5-cm, vesicles and well-defined erythematous scaly papules and plaques on the chest.