

An International, Multicenter, Randomized, Double-Blind, Parallel Group, Vehicle-Controlled, Phase 2/3 Study with Open-Label Extension Evaluating the Efficacy and Safety of Diacerein 1% Ointment for the Treatment of Generalized Epidermolysis Bullosa Simplex (EBS) (EBSshield Study/AC-203-EBS-007)

Study Overview

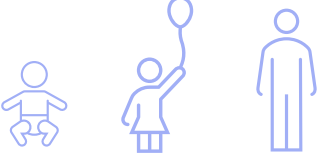




The AC-203-EBS-007 study is an international clinical trial for people with Epidermolysis Bullosa Simplex (EBS). Participants will be randomly assigned to use either diacerein 1% ointment (AC-203) or a control ointment once daily for 8 weeks, without knowing which ointment they are using. After an additional 8-week period without treatment, all participants will use diacerein 1% ointment (AC-203) for 24 weeks. Participants will report outcomes, including pain, itching, and quality of life at each visit.

Global Enrollment Number: 80-100 participants

Study Drug: Diacerein 1% ointment (AC-203) or vehicle



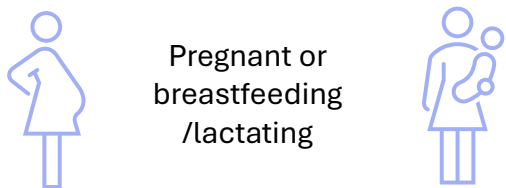
We are looking for patients with **Epidermolysis Bullosa Simplex (EBS)** to join our study.

Inclusion criteria (You must have...)

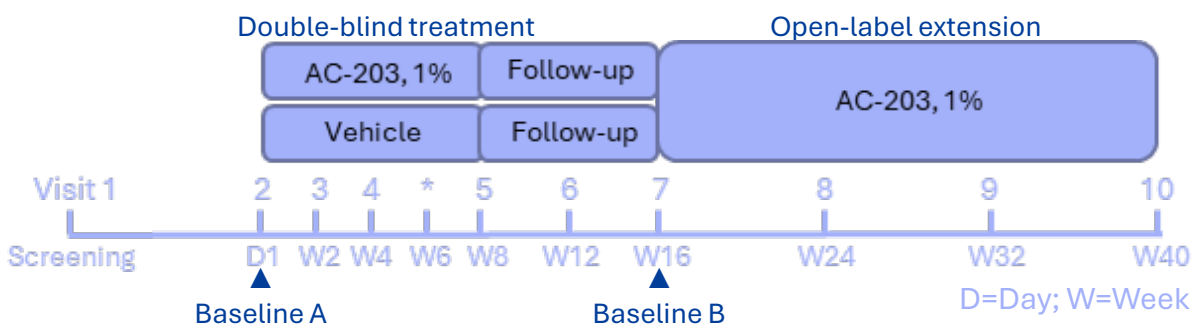
-  At least 6 months of age
-  Clinically diagnosed severe EBS or intermediate EBS
-  Autosomal dominant *KRT5* or *KRT14* gene mutation
-  EBS lesions at least 5% body surface area (BSA)
-  Treatment area IGA score at least 3


Note: If you would like to clarify whether you meet the criteria, please consult your doctor.

Exclusion criteria (You must not have...)

-  Other clinically significant skin disease(s)
-  Current cancer or a history of treatment for a cancer within 5 years
-  Pregnant or breastfeeding /lactating

Study design: You will be in this study for up to 50 weeks (screening period up to 10 weeks).



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