

## **Tralokinumab**

Resolution of: 6 January 2022  
 Entry into force on: 6 January 2022  
 Federal Gazette, BAnz AT 10 02 2022 B8

valid until: unlimited

### **Therapeutic indication (according to the marketing authorisation of 17 June 2021):**

Adtralza is indicated for the treatment of moderate-to-severe atopic dermatitis in adult patients who are candidates for systemic therapy.

### **Therapeutic indication of the resolution (resolution of 6 January 2022):**

see therapeutic indication according to marketing authorisation.

## **1. Additional benefit of the medicinal product in relation to the appropriate comparator therapy**

Adult patients with moderate-to-severe atopic dermatitis who are eligible for continuous systemic therapy

### **Appropriate comparator therapy:**

Dupilumab (in combination with topical glucocorticoids (TCS) and/or topical calcineurin inhibitors (TCI) if required)

### **Extent and probability of the additional benefit of Tralokinumab compared to Dupilumab:**

An additional benefit is not proven.

## **Summary of results for relevant clinical endpoints**

<b>Endpoint category</b>	<b>Direction of effect/ risk of bias</b>	<b>Summary</b>
Mortality	n.a.	No suitable data available.
Morbidity	n.a.	No suitable data available.
Health-related quality of life	n.a.	No suitable data available.
Side effects	n.a.	No suitable data available.
Explanations: ↑: statistically significant and relevant positive effect with low/unclear reliability of data ↓: statistically significant and relevant negative effect with low/unclear reliability of data ↑↑: statistically significant and relevant positive effect with high reliability of data ↓↓: statistically significant and relevant negative effect with high reliability of data ↔: no statistically significant or relevant difference ∅: There are no usable data for the benefit assessment. n.a.: not assessable		

## 2. Number of patients or demarcation of patient groups eligible for treatment

Adult patients with moderate-to-severe atopic dermatitis who are eligible for continuous systemic therapy

approx. 52,000 patients

## 3. Requirements for a quality-assured application

The requirements in the product information are to be taken into account. The European Medicines Agency (EMA) provides the contents of the product information (summary of product characteristics, SmPC) for Adtralza (active ingredient: tralokinumab) at the following publicly accessible link (last access: 14 December 2021):

[https://www.ema.europa.eu/en/documents/product-information/adtralza-epar-product-information\\_en.pdf](https://www.ema.europa.eu/en/documents/product-information/adtralza-epar-product-information_en.pdf)

Discontinuation of treatment should be considered for patients who do not show a response after 16 weeks of treatment. Some patients with an initial partial response may continue to benefit from fortnightly treatment continued beyond 16 weeks.

## 4. Treatment costs

### Annual treatment costs:

Adult patients with moderate-to-severe atopic dermatitis who are eligible for continuous systemic therapy

Designation of the therapy	Annual treatment costs/ patient
Medicinal product to be assessed:	
Tralokinumab	€ 8,863.47 - € 17,795.11
Appropriate comparator therapy:	
Dupilumab	€ 17,795.11

Costs after deduction of statutory rebates (LAUER-TAXE® as last revised: 1 December 2021)

Costs for additionally required SHI services: not applicable