

IMCY-0098 belongs to a new class of **active** and **specific immunotherapies** (Imotopes[™]) for the treatment of severe **autoimmune diseases**.

It has been tested previously in a first clinical study on 31 patients and demonstrated a **safe profile**.



This study will <u>test this new drug</u> for the **treatment of type 1 diabetes** in collaboration with the INNODIA European platform

Objectives of IMPACT study

- Determine the optimal treatment dose
- Test the clinical efficacy of IMCY-0098
- Confirm its safety in adults
- Detect and describe the new immune cells induced by the treatment



Conditions of participation

- Age ≥18 and <45 years at consent
- Diagnosis of T1D within 9 weeks at screening (date of 1st insulin injection)
- Being on insulin treatment
- Having at least one diabetes-related autoantibody present at screening
- Having random C-peptide levels ≥200 pmol/L measured at screening
- Being HLA DR4 positive
 > Up to 24 DR4 negative (but DR3 positive) patients will be able to participate in a substudy

Study visits and main assessments



Study program outline

- 7 treatment visits and 3 follow-up visits
- 2 injections of IMCY-0098 or placebo at each treatment visit
- Total study participation will last approximately 1 year including screening

In addition to the tests and procedures used to check your health (measurement of BP, pulse, physical exam. etc), the following specific tests will be performed:

- Mixed Meal Tolerance Test (MMTT)
- Blood samples for immune analysis
- Follow-up of potential injection site reactions
- Dry Blood Spot (DBS) at home

Site contact information