

Participant travel reimbursement enhancement for clinical trials



An overview of Markel's new travel cover enhancement designed to support clinical trials.

The enhancement can provide trial sponsors reimbursement for eligible expenses associated with international travel by approved participants to and from clinical-trial sites, subject to policy terms and conditions. Participants are not insured under the policy and do not receive insurance payments from Markel. Insurance benefits are payable to the trial sponsor by way of reimbursement, subject to the policy terms and conditions.



HELPING REMOVE BARRIERS TO PARTICIPATION

Reimbursement of eligible travel-related expenses to trial sponsors is designed to help them address some of the financial barriers associated with participant travel to specialist trial sites.



SUPPORT THROUGHOUT THE JOURNEY

Dedicated 24-hour travel assistance is available from global healthcare services provider, Healix, providing participants with medical assistance and the coordination of appropriate care while travelling.



GLOBAL AVAILABILITY

Accessible through brokers worldwide, the enhancement supports clinical trials involving participants travelling across international borders.



GET IN TOUCH WITH OUR TEAM FOR MORE INFORMATION



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Important

- All cover, benefits and services are subject to policy terms and conditions.
- Participants must be registered under the policy and medically assessed as fit to travel by their doctor.
- The insurance does not determine whether someone is eligible to participate in a clinical trial or replace clinical assessment and informed consent processes.
- Eligibility to participate remains subject to the trial sponsor's protocols, investigator assessment, informed consent requirements and applicable regulatory requirements.

