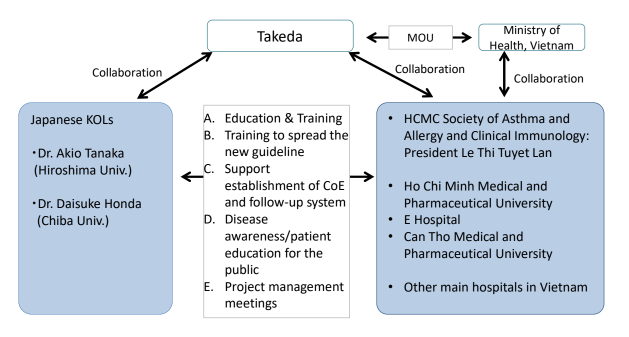
Project on establishing diagnostic methods, strengthening treatment, and developing guidelines for hereditary angioedema in Vietnam

[BACKGROUND]

- Disease awareness of HAE in Vietnam is extremely low, testing and diagnostic techniques have not been established, and there are no core centers of care nor treatment guidelines in place.
- The Vietnamese Ministry of Health, academic societies, and physicians recognize the necessity to improve the environment for HAE.

[PROJECT OVERVIEW]

- Year 1 (2021): Raise awareness of the disease and strengthen the capacity of Vietnamese healthcare professionals and begin formulating local HAE treatment guidelines.
- Year 2 (2022): Establish testing and diagnostic techniques, formulate local HAE treatment guidelines, raise awareness of the disease among Vietnamese citizens, and begin discussions on establishing Center of Excellence (CoE) and follow-up system.
- Year 3 (2023): Establish CoE and follow-up system, establish a collaborative system among medical institutions, improve access to medicines and treatment rates by obtaining marketing approval for new HAE medicines and getting them on insurance coverage, and create an Asian HAE/rare disease treatment ecosystem centered on Japan through expansion to other regions.



July-December 2023 Education and Training (Testing, Diagnosys, and Treatment) (On-site or Web)

B. August-October 2023 Dissemination training on the new guideline (Web)

C. November 2023 Support for CoE establishment and follow-up system development (On-site or web)

D. July-August 2023 Disease awareness/patient education for the public (Web)

E. July 2023-January 2024 Project management meeting (Web)