Tay’s syndrome, a rare autosomal recessive disorder, characterized by ichthyosiform erythroderma, trichothiodystrophy, brittle hair and nails, intellectual impairment, decreased fertility, short stature, progeria-like facies and photosensitivity [1-3]. There is no specific treatment for this genetic disorder. However, some authors reported control of ichthyosis in a case of Tay’s syndrome by topical application of tazarotene [3].

This syndrome is reported first by, Dr. Tay Chong Hai in Singapore in the year 1969 [4]. Dr Tay reported it in 2 brothers and a sister, with first-cousin parents of Chinese extraction. One of the children had hypogammaglobulinemia, and one died at age 2 months of intestinal obstruction. Erythroderma was particularly striking at birth.

Chong Hai Tay, was born in 1932 (Fig. 1). He is the first Singapore physician to have a disease named for him in the Western medical literature [5]. He authored more than hundred scientific articles. He wrote also poems. He established the National Arthritis Foundation in 1984, and was chairman for 14 years.

He is currently in private practice and a consultant physician and rheumatologist at Mt. Elizabeth Hospital, Gleneagles Hospital, Mt. Alvernia Hospital and East Shore Hospital. Dr Tay was the first editor of The Scientific Victorian [5]. Among his many other scientific contributions, he also, reported ten patients who were presenting with acute polyarthritis and hypereosinophilia of unknown causation [6].

REFERENCES