

Further Halotyrosine Derivatives from the Marine Sponge *Suberea* aff. *praetensa*[§]

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Z. Naturforsch. **57c**, 732–738 (2002); received April 9/May 21, 2002

Suberea aff. *praetensa*, Halotyrosine Derivatives, Anticancer Activities

Reexamination of the marine sponge *Suberea* aff. *praetensa*, (Row) from the Gulf of Thailand furnished in addition to bromotyrosine derivatives found previously 5-bromo- and 5-chlorocavernicolin, cavernicolins 1 and 2, two other brominated tyrosine metabolites, a known bisoxazolidone and a new unusual rearranged tyrosine metabolite subereatensin. Several of the metabolites exhibited significant inhibitory effects against five human cancer cell lines.