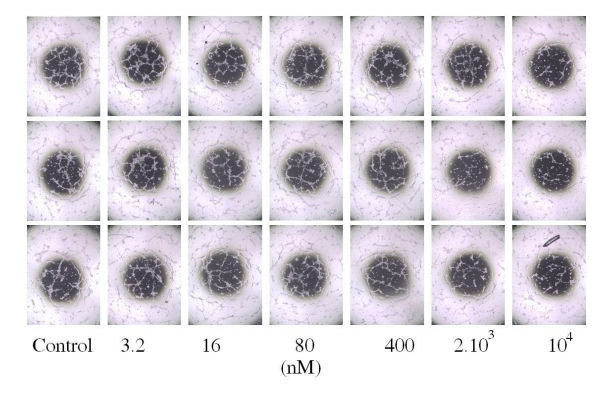
A low molecular weight protein from the Sea Anemone Anemonia viridis with an anti Angiogenic activity.



Supplementary results

Figure 1S. HMEC tubulogenesis assay with the 28' fraction with concentrations going from 10 μ M to 3.2 nM in a triplicate experiment. A network with interconnected lines similar to control was observed only at 3.2 nM.

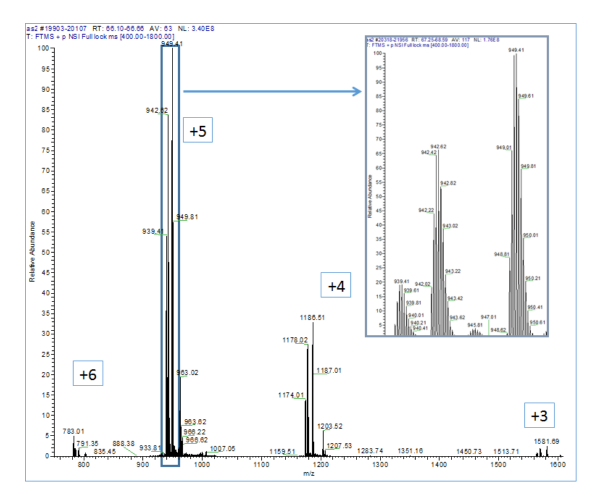


Fig 2S. High resolution nano-HPLC MS of fraction 28'. The numbers in square indicate the charge state of the peptides, the insert shows the isotopic envelopes of the quintupled charged main species

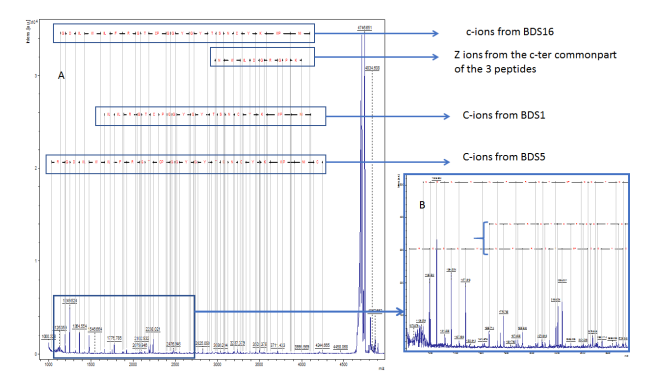


Fig 3S. Maldi ISD of fraction 28. A:annotation of the ions obtained by Maldi ISD of the three main species of the fraction. B: The insert shows the common n-terminal part of BDS1 and BDS5.

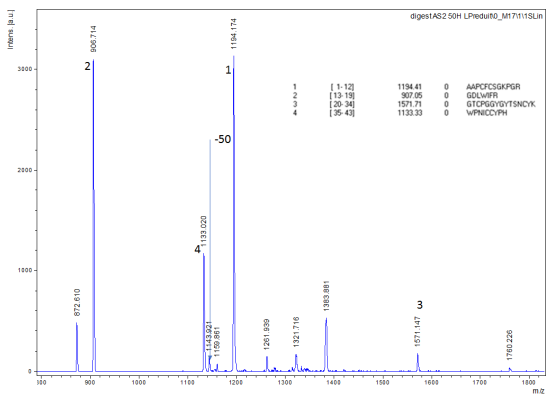


Fig 4S. MALDI spectra of digest of the 28' fraction after reduction before Nano-HPLC ms/ms analysis on Orbitrap Q-exactive: The peaks 1 to 4 match with the sequence of BDS-5(table in insert), A is specific to BDS1 with a Leu/PHE mutation, B match with the BDS16 sequence and was de Novo sequenced by hplc-ms/ms as shown below.

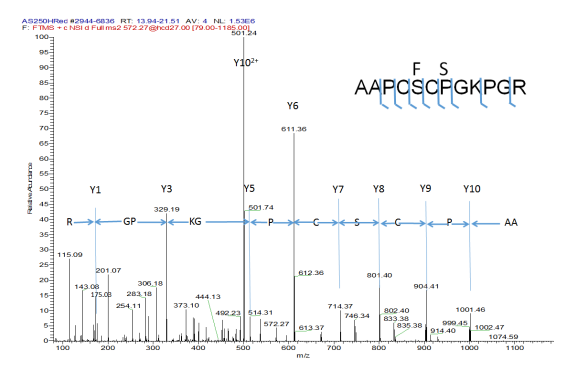


Fig 5S.MS/MS spectrum of the peptide of the N-terminal part of BDS-16 with two mutations (S5et P7 instead of F5 et S7 for BDS1 and BDS5). The spectrum is annotated by the y-ions generated by the HCD fragmentation from the Q-exactive Orbitrap.

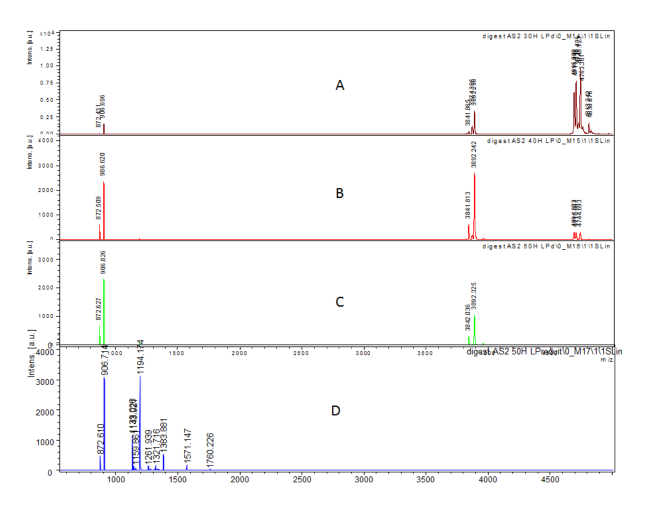


Fig 6S.Digestion of the non-reduced fraction from 30 to 70 hours. A: 30 H, B: 40 H C: 70 H D: Reduction of the digest 70H