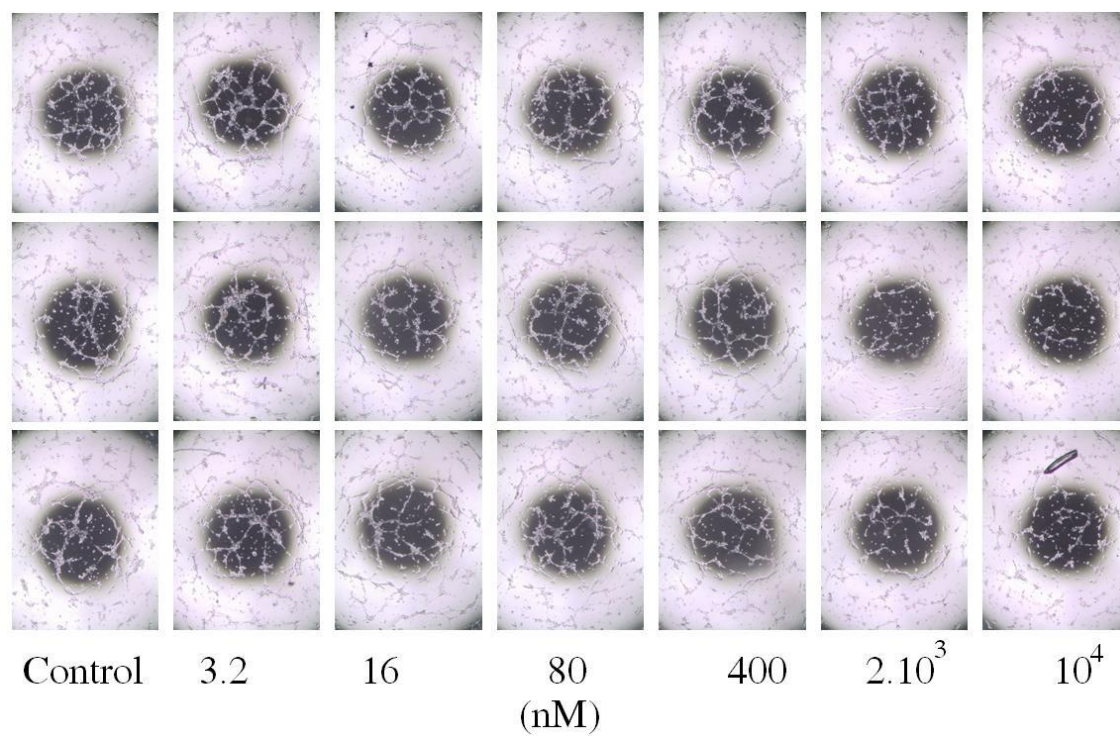
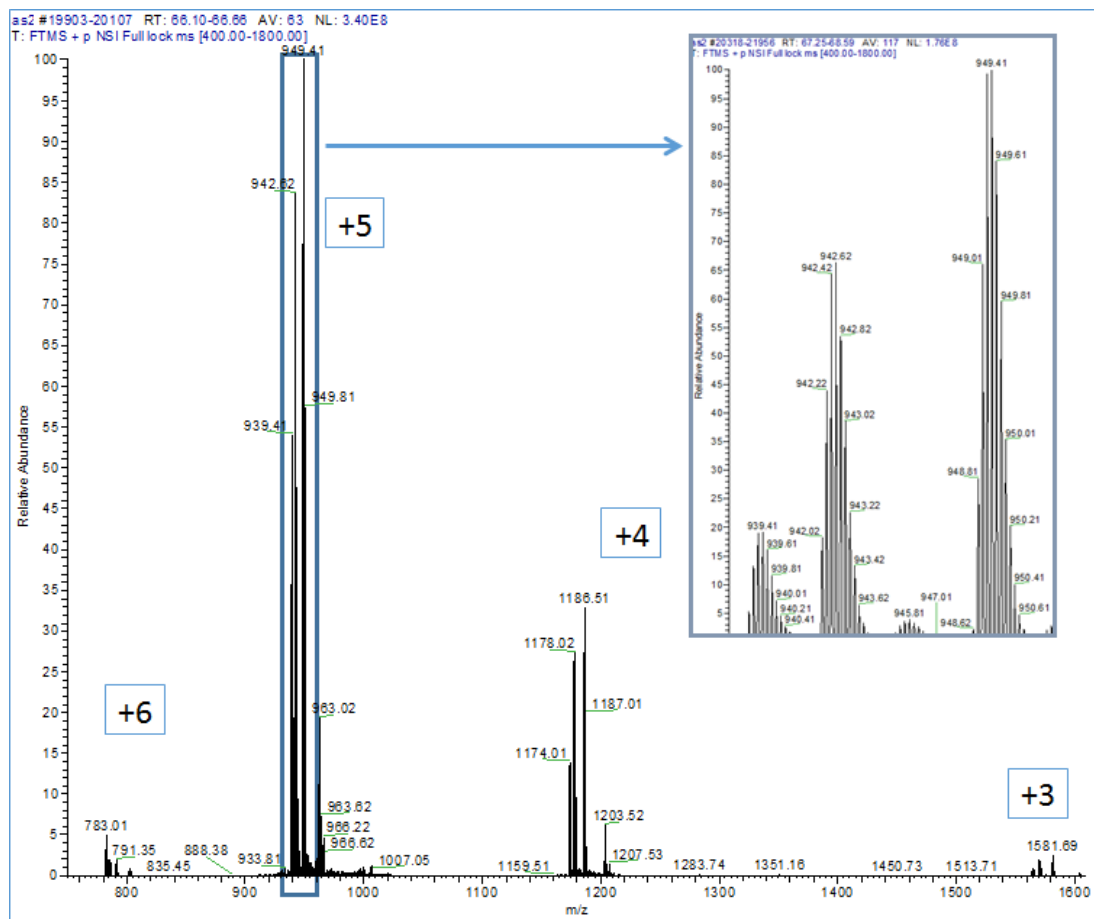


**A low molecular weight protein from the Sea Anemone  
*Anemonia viridis* with an anti Angiogenic activity.**

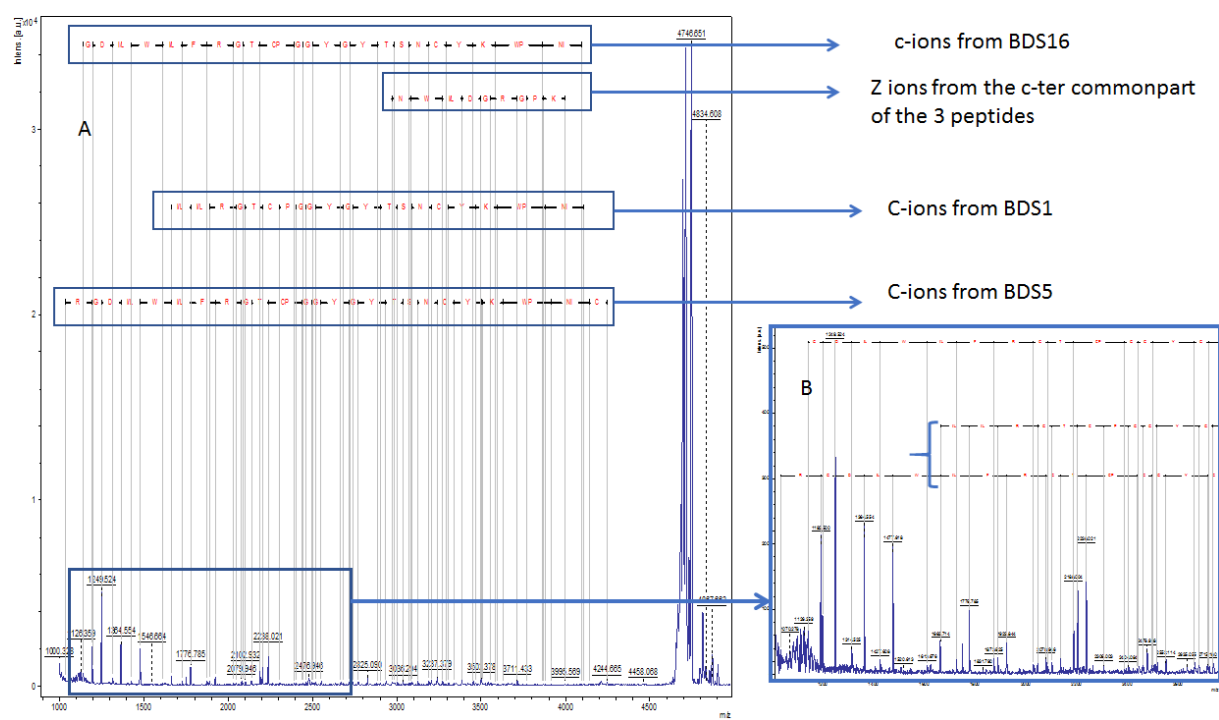
**Supplementary results**



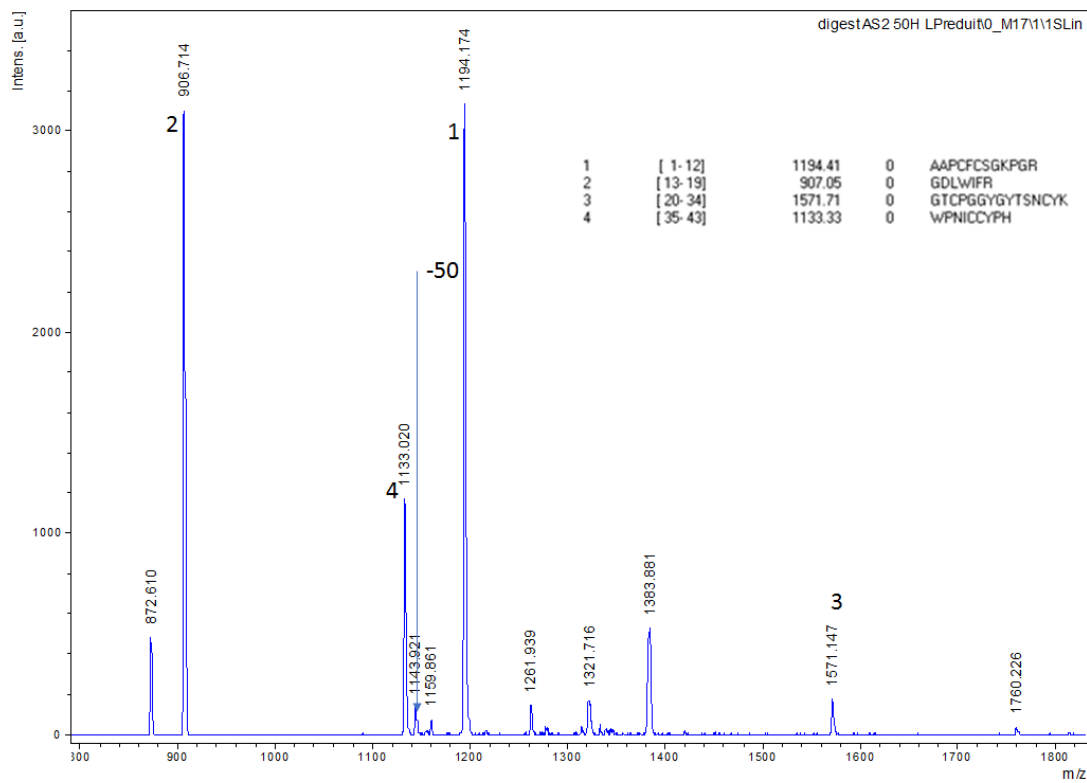
**Figure 1S.** HMEC tubulogenesis assay with the 28' fraction with concentrations going from 10  $\mu$ M to 3.2 nM in a triplicate experiment. A network with interconnected lines similar to control was observed only at 3.2 nM.



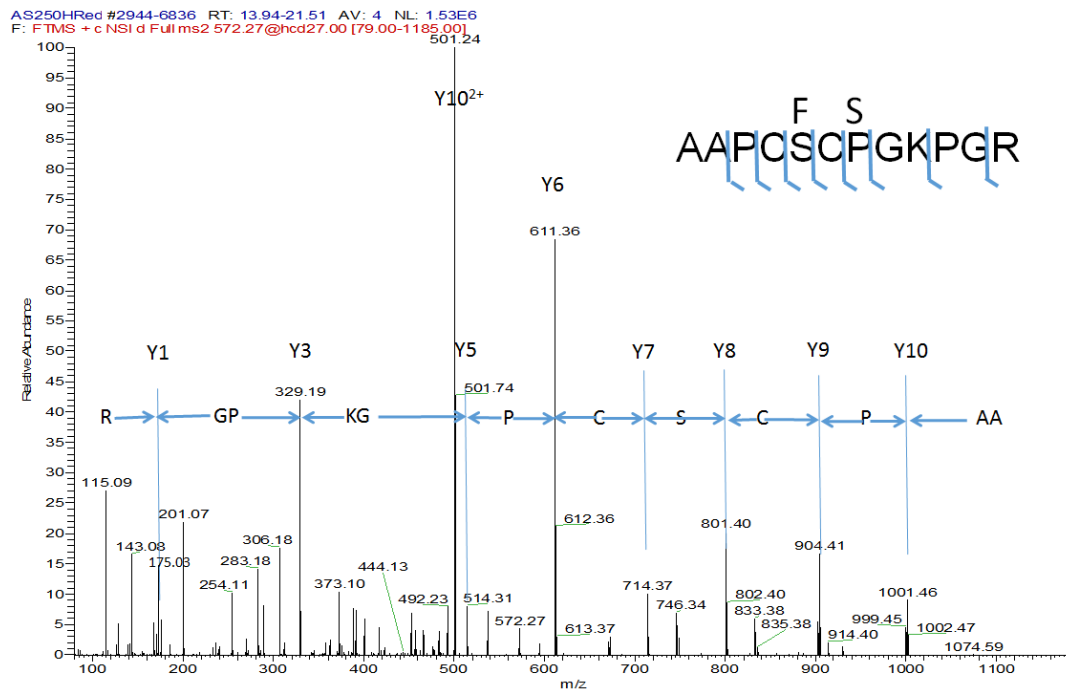
**Fig 2S.** High resolution nano-HPLC MS of fraction 28'. The numbers in square indicate the charge state of the peptides, the insert shows the isotopic envelopes of the quintupled charged main species



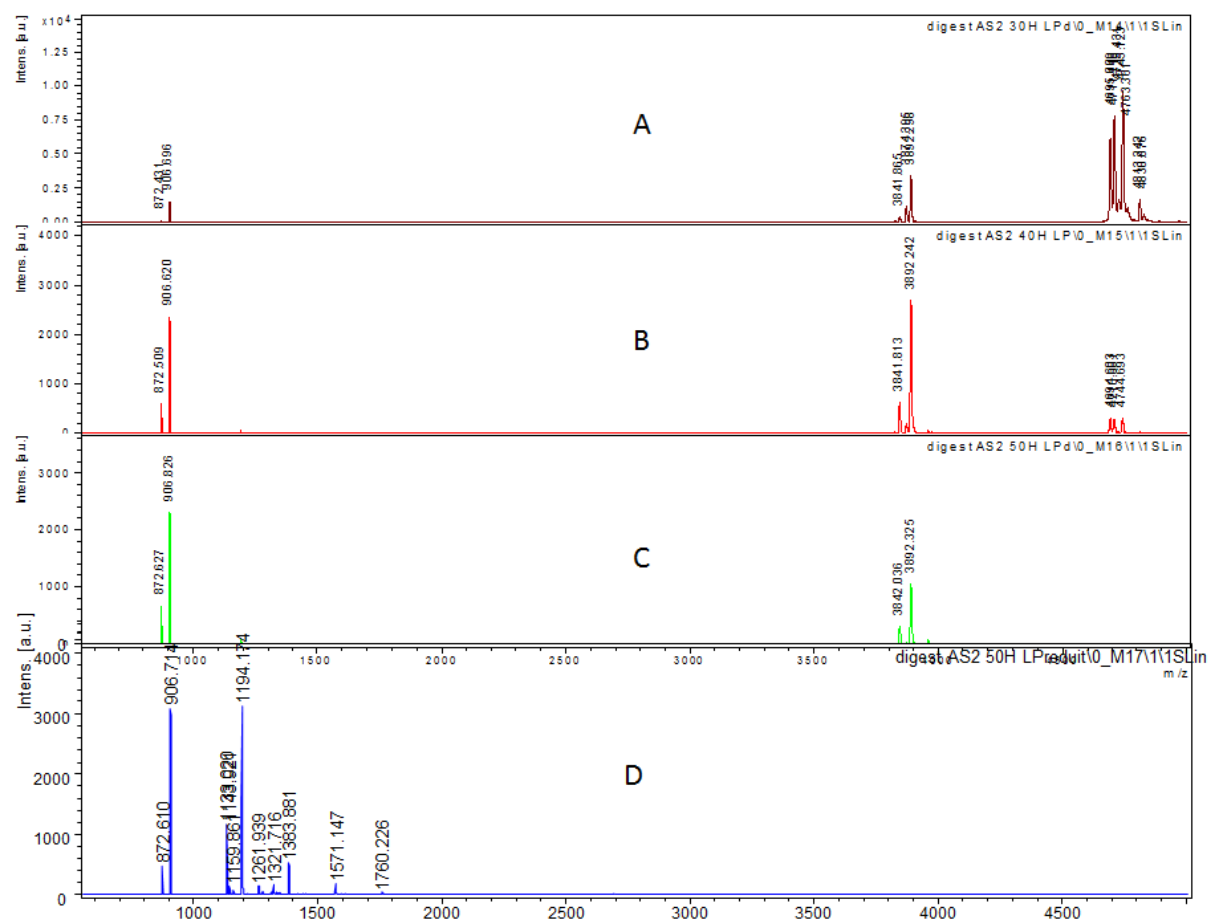
**Fig 3S.** Maldi ISD of fraction 28. A:annotation of the ions obtained by Maldi ISD of the three main species of the fraction. B: The insert shows the common n-terminal part of BDS1 and BDS5.



**Fig 4S.** MALDI spectra of digest of the 28<sup>3</sup> 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