Metalloproteinases diversity in the venom gland of Peruvian spider Loxosceles laeta revealed by transcriptome analysis

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Abstract

Envenomation caused by spiders from Loxosceles genus (brown spiders) is a worldwide public health problem. Loxosceles their venom is composed of several toxins responsible for dermonecrotic, hemorrhagic and edema effects. In Peru, L. laeta is considered the most medical relevant species. A family of metalloproteases, also named astacin-like proteins, was described in Loxosceles venom with great importance for hemostatic disorders in natural or experimental envenomations. A new generation sequencing library of venom extracted from the Peruvian spider, L. laeta, was constructed for the first time using the TruSeq TM RNA Sample Prep Kit v3 Set A (Illumina) kit and the sequencing was performed on the MiSeq by the paired-end technique for identification of molecular diversity of metalloproteases toxins. In this work, we describe some of the identified metalloproteases enzymes with a high degree of identity (over 50%) with molecules from other Loxosceles spp spiders. Results obtained in this work represent the first landscape of components of a Peruvian spider venom gland, revealing the complexity of molecules expressed in this tissue, with great potential for future uses in medical and evolutionary studies.

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