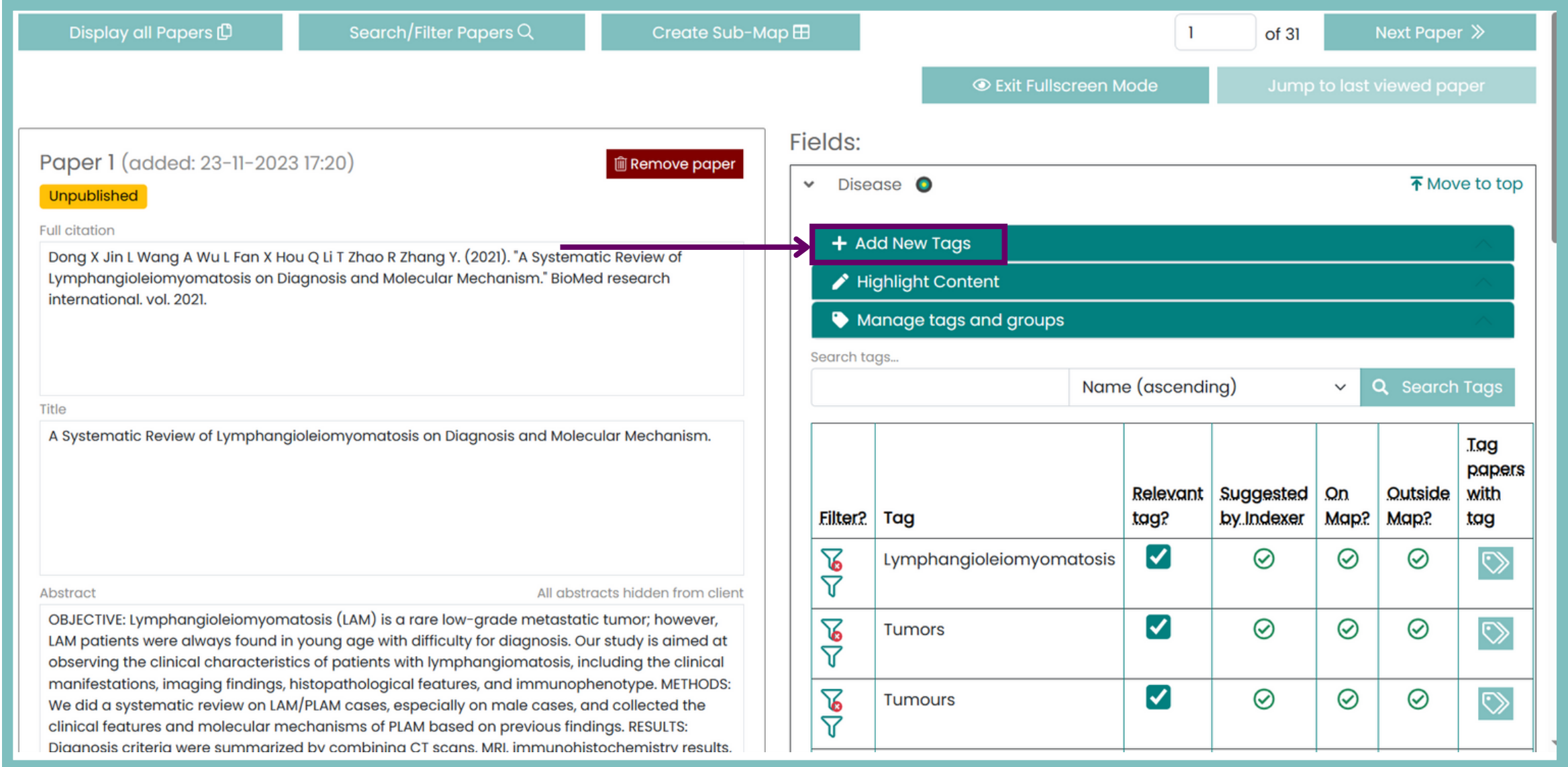


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Dong X Jin L Wang A Wu L Fan X Hou Q Li T Zhao R Zhang Y. (2021). "A Systematic Review of Lymphangioleiomyomatosis on Diagnosis and Molecular Mechanism." BioMed research international. vol. 2021.

Title
A Systematic Review of Lymphangioleiomyomatosis on Diagnosis and Molecular Mechanism.

Abstract All abstracts hidden from client
OBJECTIVE: Lymphangioleiomyomatosis (LAM) is a rare low-grade metastatic tumor; however, LAM patients were always found in young age with difficulty for diagnosis. Our study is aimed at observing the clinical characteristics of patients with lymphangiomas, including the clinical manifestations, imaging findings, histopathological features, and immunophenotype. METHODS: We did a systematic review on LAM/PLAM cases, especially on male cases, and collected the clinical features and molecular mechanisms of PLAM based on previous findings. RESULTS: Diaanosis criteria were summarized by combinina CT scans. MRI. immunohistochemistrv results.

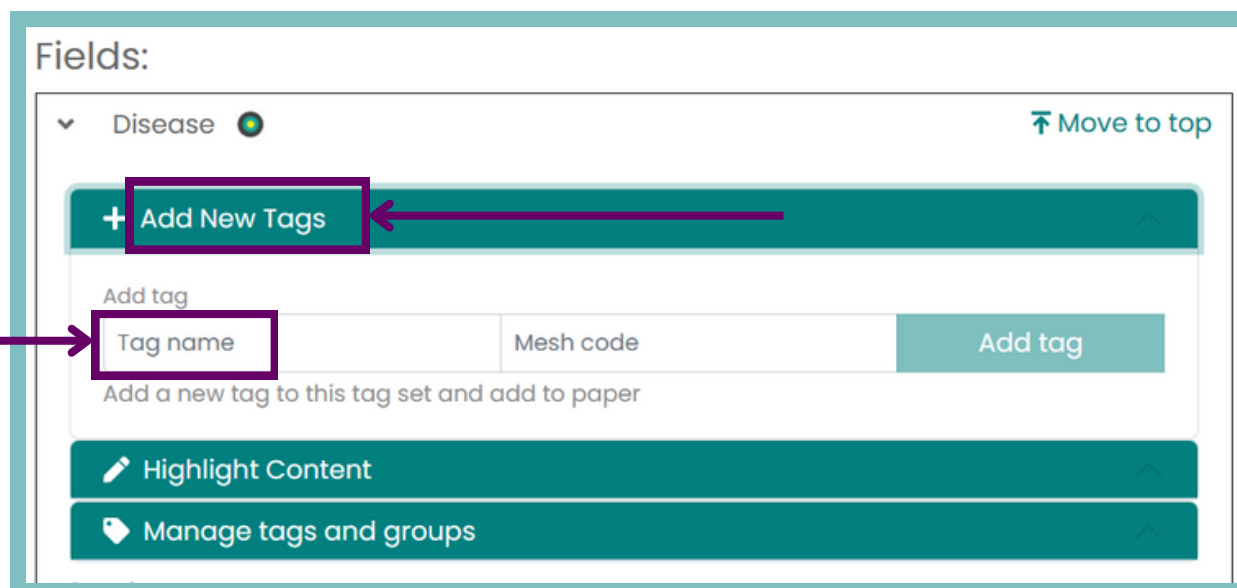
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