

POSTER PRESENTATION

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The lebercillin-like protein is embedded in a ciliary protein network and is preferentially expressed in motile cilia

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Mutations in *LCA5* are causative for Leber congenital amaurosis, a severe hereditary retinal dystrophy in humans. Lebercillin, encoded by *LCA5*, localizes to connecting cilia of photoreceptor cells in the retina and specifically interacts with the intraflagellar transport (IFT) machinery. Bioinformatic analysis has identified lebercillin-like protein, previously known as C21orf13, as a lebercillin homolog in humans. In this study, we have characterized the molecular properties of lebercillin-like protein by defining the lebercillin-like interactome and assessing its (sub)cellular localization in ciliated cells. We show that lebercillin-like protein is embedded in a ciliary protein network and specifically localizes at the basal body and ciliary axoneme of ciliated cells, like lebercillin. mRNA expression studies indicate that lebercillin-like protein is preferentially expressed in tissues featuring motile cilia and/or flagella. Based on these data and bioinformatic co-expression profiling, we suggest that *LCA5L* is a likely candidate gene for motile ciliopathies such as Primary Ciliary Dyskinesia (PCD).

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