

Vivet Therapeutics raises €37.5 million in Series A financing

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Vivet Therapeutics is an emerging biotechnology company developing novel gene therapies for rare, inherited metabolic diseases. The company has raised €37.5 million in a Series A financing round that involved a syndicate of leading international life sciences investors led by Novartis Venture Fund and Columbus Venture Partners, and including Roche Venture Fund, HealthCap, Kurma Partners and Ysios Capital.

The funds will be used by Vivet to advance a diversified pipeline of gene therapy programs targeting rare, inherited metabolic diseases, including Wilson Disease, progressive familial intrahepatic cholestasis type 2 (PFIC2), progressive familial intrahepatic cholestasis type 3 (PFIC3) and citrullinemia type I.

Vivet is building its pipeline based on novel technology developed through its partnerships with, and exclusive licenses from, the Fundación para la Investigación Médica Aplicada (FIMA), a not-for-profit foundation at the Centro de Investigación Médica Aplicada (CIMA), University of Navarra based in Pamplona, Spain; and Massachusetts Eye and Ear (MEE), Boston, USA. The licenses cover exclusive use of novel proprietary AAV vector gene therapy technology to treat metabolic diseases and certain MEE patent-protected Anc80 AAV gene therapy vectors. Anc80 is a next-generation gene therapy technology designed to increase gene expression levels in the liver, while reducing the risk of undesired immunogenicity.