

ERYTECH announces three data presentations at the 2015 AACR Annual Meeting

Lyon (France), March 24th 2015 – ERYTECH Pharma (Euronext Paris: ERYP & OTC US: EYRY), the French biopharmaceutical company that develops innovative ‘tumor starvation’ treatments for acute leukemia and other oncology indications with unmet medical needs, announces the upcoming presentation of three abstracts, at the American Association for Cancer Research (AACR) taking place in Philadelphia, PA, USA from 18 to 22 April 2015.

Details for the three posters are as follows:

- A phase III, multicenter, randomized study evaluating the efficacy and safety of erythrocyte encapsulated L-asparaginase (ERY001) versus native L-asparaginase (L-asp) in combination with COOPRALL regimen with first relapse of acute lymphoblastic leukemia
Abstract # CT113 - Sunday, April 19, 2015, 1:00PM 5:00PM - Location: section 22
- Low asparagine synthetase expression and in vitro sensitivity highlights L-asparaginase potential for the treatment of aggressive lymphomas
Abstract # 5369 – Wednesday, April 22, 2015, 8:00AM-12:00PM - Location: section 28
- Methioninase loaded erythrocytes: a promising drug for L-methionine restriction therapy in cancer
Abstract # 5330 – Wednesday, April 22, 2015, 8:00AM-12:00PM - Location: section 27

The meeting abstracts can be viewed online through the AACR website at www.aacr.org

About ERYTECH and ERY-ASP/GRASPA®: www.erytech.com

Created in Lyon in 2004, ERYTECH is a French biopharmaceutical company providing new prospects for cancer patients, particularly those with acute leukemia and selected solid tumors.

By encapsulating the asparaginase enzyme in red blood cells, ERYTECH has developed ERY-ASP/GRASPA®, an original treatment that targets cancer cells through “tumor starvation” while significantly reducing the side effects for patients. ERY-ASP/GRASPA® has recently announced positive Phase III data in Acute Lymphoblastic Leukemia (ALL) and is in Phase IIb clinical trial in Acute Myeloid Leukemia (AML) in Europe. The product is also in Phase I/II clinical development in ALL in the USA.

Every year about 50,000 patients are diagnosed with Acute Lymphoblastic Leukemia (ALL) or Acute Myeloid Leukemia (AML), the two forms of acute leukemia. Today, for about 80% of these patients, mainly adults and relapsing patients, current forms of asparaginase cannot be used due to their toxicity. With a presumed improved safety profile, ERY-ASP/GRASPA® is being developed to allow all leukemia patients to be treated, even the most fragile ones, representing a market opportunity of more than EUR 1 billion.

The company is also developing other indications in solid tumors and certain orphan indications outside oncology. A Phase II study in pancreas cancer is ongoing and the company is exploring other solid tumor indications for ERY-ASP.

ERYTECH has obtained orphan drug designations for ERY-ASP/GRASPA® in ALL, AML and pancreas cancer, both in Europe and the USA, and has its own GMP-approved and operational manufacturing site in Lyon (France), and a site for clinical production in Philadelphia (USA).

The company has concluded licensing and distribution partnership agreements for ALL and AML in Europe with Orphan Europe (Recordati Group), and for ALL with TEVA in Israel.

ERYTECH is listed on Euronext regulated market in Paris (ISIN code: FR0011471135, ticker: ERYP) and is part of the CAC Healthcare, CAC Pharma & Bio, CAC Mid & Small, CAC All Tradable, EnterNext PEA-PME 150 and Next Biotech indexes. ERYTECH is also listed in the US under an ADR level 1 program (OTC, ticker EYRY).

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Ticker : EYRY

Forward-looking information

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