Edinburgh, United Kingdom, September 12, 2017 (GLOBE NEWSWIRE) – NuCana plc, a clinical-stage biopharmaceutical company developing a portfolio of novel anti-cancer medicines called ProTides™, announced the presentation of data from its recently completed trial of its lead product candidate, NUC-1031 (Acelarin®), at the European Society for Medical Oncology (ESMO) 2017 Congress held September 8th-12th, 2017 in Madrid, Spain. Results from NuCana’s Phase 1b trial showed that Acelarin, when combined with carboplatin, was well tolerated and demonstrated clinical activity in women with recurrent platinum-resistant and platinum-sensitive ovarian cancer. An overall response rate of 39% was observed amongst the 23 evaluable patients, including 1 (4%) who achieved a complete response, 8 (35%) with partial responses, and 13 (57%) with stable disease that lasted at least 12 weeks. This yielded an overall disease control rate of 96% (22 patients). The responses were durable, with an average progression free survival of 7.4 months. The most common adverse events across all dose levels were neutropaenia, leukopaenia and thrombocytopaenia. No unexpected adverse events were observed with the combination to date.

All patients in the study were previously treated with an average of three prior chemotherapy regimens. Seventeen of the evaluable patients were either refractory or resistant to their last platinum-containing regimen.

“The fact that the Acelarin combination with carboplatin achieved these results in heavily pre-treated and platinum-resistant patients clearly demonstrates Acelarin is a very active agent,” said Dr. Sarah Blagden, Associate Professor of Experimental Cancer Medicine at the University of Oxford and Chief Investigator of the Phase 1b study. Professor Blagden added, “importantly, the favorable toxicity profile of Acelarin enabled us to combine it with carboplatin at AUC 5, whereas with gemcitabine, carboplatin has to be given at AUC 4. Thus, we are able to deliver both Acelarin and carboplatin at their optimal dose.”

Hugh Griffith, NuCana’s Chief Executive Officer, said: “The high disease control rate and durable responses achieved with the combination of Acelarin and carboplatin are exciting. We remain focused on advancing Acelarin’s development for the treatment of patients with ovarian cancer as well as exploring its use for the treatment of other solid tumours.”

About NuCana
NuCana® is a clinical-stage biopharmaceutical company focused on significantly improving treatment outcomes for cancer patients by applying our ProTide™ technology to transform some of the most widely prescribed chemotherapy agents, nucleoside analogs, into more effective and safer medicines. While these conventional agents remain part of the standard of care for the treatment of many solid tumours, their efficacy is limited by cancer cell resistance mechanisms and they are often poorly tolerated. Utilising our
proprietary technology, we are developing new medicines, ProTides, designed to overcome key cancer resistance mechanisms and generate much higher concentrations of anti-cancer metabolites in cancer cells. Our most advanced ProTide candidates, Acelarin® and NUC-3373, are new chemical entities derived from the nucleoside analogs gemcitabine and 5-fluorouracil, respectively, two widely used chemotherapy agents. Acelarin is currently being evaluated in four clinical studies across several solid tumour indications, including ovarian cancer, biliary cancer and pancreatic cancer. NUC-3373 is currently in a Phase 1 study for the potential treatment of a wide range of advanced solid tumour cancers. For more information, please visit: www.nucana.com.

Forward-Looking Statements
This press release may contain “forward-looking” statements that are based on the beliefs and assumptions and on information currently available to management of NuCana plc (the “Company”). All statements other than statements of historical fact contained in this press release are forward-looking statements, including statements concerning the initiation, timing, progress and results of clinical trials of the Company’s product candidates. In some cases, you can identify forward-looking statements by terminology such as “may,” “will,” “should,” “expects,” “plans,” “anticipates,” “believes,” “estimates,” “predicts,” “potential” or “continue” or the negative of these terms or other comparable terminology. Forward-looking statements involve known and unknown risks, uncertainties and other factors that may cause the Company’s actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements. Forward-looking statements represent the Company’s beliefs and assumptions only as of the date of this press release. Although the Company believes that the expectations reflected in the forward-looking statements are reasonable, it cannot guarantee future results, levels of activity, performance or achievements. Except as required by law, the Company assumes no obligation to publicly update any forward-looking statements for any reason after the date of this press release to conform any of the forward-looking statements to actual results or to changes in its expectations.

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