



CryoLife Announces Release Date and Teleconference Call Details for 2011 Third Quarter Financial Results

October 17, 2011

ATLANTA, Oct. 17, 2011 /PRNewswire via COMTEX/ -- CryoLife, Inc. (NYSE: CRY), a leading tissue processing and medical device Company focused on cardiac and vascular surgery, announced today that 2011 third quarter financial results will be released on Thursday, October 27, 2011. On that day, the Company will hold a teleconference call and live webcast at 10:00 a.m. Eastern Time to discuss the results, followed by a question and answer session hosted by Steven G. Anderson, president and chief executive officer of CryoLife, Inc.

To listen to the live teleconference, please dial 201-689-8433 a few minutes prior to 10:00 a.m. A replay of the teleconference will be available October 27 through November 3 and can be accessed by calling (toll free) 877-660-6853 or 201-612-7415. The account number for the replay is 244 and the conference number is 380521.

The live webcast and replay can be accessed by going to the Investor Relations section of the CryoLife web site at <http://www.cryolife.com/> and selecting the heading Webcasts & Presentations.

About CryoLife, Inc.

Founded in 1984, CryoLife, Inc. is a leader in the processing and distribution of implantable living human tissues for use in cardiac and vascular surgeries throughout the U.S. and Canada. CryoLife's CryoValve® SG pulmonary heart valve, processed using CryoLife's proprietary SynerGraft® technology, has FDA 510(k) clearance for the replacement of diseased, damaged, malformed, or malfunctioning native or prosthetic pulmonary valves. CryoLife's CryoPatch® SG pulmonary cardiac patch has FDA 510(k) clearance for the repair or reconstruction of the right ventricular outflow tract (RVOT), which is a surgery commonly performed in children with congenital heart defects, such as Tetralogy of Fallot, Truncus Arteriosus, and Pulmonary Atresia. CryoPatch SG is distributed in three anatomic configurations: pulmonary hemi-artery, pulmonary trunk, and pulmonary branch. CryoLife's BioGlue® Surgical Adhesive is FDA approved as an adjunct to sutures and staples for use in adult patients in open surgical repair of large vessels. BioGlue is also CE marked in the European Community and approved in Canada and Australia for use in soft tissue repair and was recently approved in Japan for use in the repair of aortic dissections. CryoLife's BioFoam(TM) Surgical Matrix is CE marked in the European Community for use as an adjunct in the sealing of abdominal parenchymal tissues (liver and spleen) when cessation of bleeding by ligature or other conventional methods is ineffective or impractical. CryoLife distributes PerClot®, an absorbable powder hemostat, in the European Community. CryoLife, through its subsidiary Cardiogenesis Corporation, specializes in the treatment of cardiovascular disease and devices that treat severe angina. Its market leading FDA-approved Holmium: YAG laser system and single use fiber-optic delivery systems are used to perform a surgical procedure known as Transmyocardial Revascularization (TMR).

For additional information about the company, visit CryoLife's Web site: <http://www.cryolife.com/>.

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SOURCE CryoLife, Inc.