

New Influenza Test Offers Labs Advanced Performance and Reliable Results

The Associated Press

BD Diagnostics, a segment of BD (Becton, Dickinson and Company) (NYSE: BDX), a leading global medical technology company, announced today that it received 510(k) clearance from the U.S. Food and Drug Administration (FDA) for nasopharyngeal wash, aspirate and swab in transport media specimens on the BD VeritorT System for Rapid Detection of Flu A+B. This new product is cleared for use in clinical settings.

The new BD Veritor System for Rapid Detection of Flu A+B laboratory kit is specifically configured for testing liquid specimens obtained via nasopharyngeal wash, aspirate or swab in transport media. The system, with proprietary technologies, eliminates the subjective result interpretation of visually read assays and helps deliver an accurate read by providing reliable, objective results on a hand held reader with an easy-to-read digital display. The BD Veritor System for Rapid Detection of Flu A + B has demonstrated proven performance versus polymerase chain reaction tests (PCR). This is the only rapid diagnostic flu test that has been referenced and FDA cleared against this high-sensitivity standard.

"Visually read rapid flu tests have shown variable performance results and are prone to errors when interpreting results," said Tom Polen, President, BD Diagnostics - Diagnostic Systems. "The BD Veritor System provides an advanced rapid diagnostic test for influenza A+B by offering accuracy, consistency, and convenience along with an objective read that laboratorians can trust." The BD Veritor System utilizes Advanced Particle and Adaptive Read Technologies coupled with a special hand-held reader. The Advanced Particle Technology along with improved chemistries helps increase the sensitivity of the test. The Adaptive Read Technology helps reduce false-positive results by examining and compensating for many of the effects of non-specific binding which improves specificity.

The clinical lab version joins the previously FDA-cleared and CLIA- waived BD Veritor System for Rapid Detection of Flu A+B. This assay for Rapid Detection of Flu A+B on the BD Veritor System represents the first of many planned assays on this new platform.

For more information on the BD Veritor System, please visit www.bd.com/ds/veritorsystem.

About BD BD is a leading global medical technology company that develops, manufactures and sells medical devices, instrument systems and reagents. The Company is dedicated to improving people's health throughout the world. BD is focused on improving drug delivery, enhancing the quality and speed of diagnosing infectious diseases and cancers, and advancing research, discovery and production

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Published on Medical Design Technology (<http://www.mdtmag.com>)

of new drugs and vaccines. BD's capabilities are instrumental in combating many of the world's most pressing diseases. Founded in 1897 and headquartered in Franklin Lakes, New Jersey, BD employs approximately 29,000 associates in more than 50 countries throughout the world. The Company serves healthcare institutions, life science researchers, clinical laboratories, the pharmaceutical industry and the general public. For more information, please visit www.bd.com.

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SOURCE BD-Becton Dickinson -0- 09/20/2012 /Web Site: <http://www.bd.com>
(NYSE:BDX) / CO: BD-Becton Dickinson; BD Diagnostics ST: New Jersey IN: HEA MTC
SU: PDT PRN -- NY77296 -- 0000 09/20/2012 12:00:00 EDT <http://www.prnewswire.c>

Source URL (retrieved on 01/26/2015 - 6:19am):

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