

DECLARATION OF CONFORMITY

Diamond Diagnostics, Inc. hereby ensures and declares that the product(s) listed below comply with the requirement of In Vitro Diagnostics Medical Devices Regulation EU 2017/746 and Directive 2011/65/EU.

Diamond Diagnostics Inc. ezúton biztosítja és kijelenti, hogy az alább felsorolt termék(ek) megfelelnek az In Vitro Diagnostics Medical Devices EU 2017/46 rendelet és a 2011/65/EU irányelv követelményeinek.

Diamond Diagnostics Inc. versichert und erklärt hiermit, dass das/die unten aufgeführte(n) Produkt(e) den Anforderungen der In-vitro-Diagnostik-Medizinprodukte-Verordnung EU 2017/46 und der Richtlinie 2011/65/EU entsprechen.

Diamond Diagnostics Inc. garantit et déclare par la présente que le ou les produits répertoriés ci-dessous sont conformes aux exigences du règlement sur les dispositifs médicaux de diagnostic in vitro UE 2017/46 et de la directive 2011/65/EU.

Diamond Diagnostics Inc. por la presente garantiza y declara que los productos enumerados a continuación cumplen con los requisitos del Reglamento de dispositivos médicos de diagnóstico in vitro EU 2017/46 y la Directiva 2011/65/EU.

Diamond Diagnostics Inc. 特此确保并声明下列产品符合体外诊断医疗器械法规 EU 2017/46 和指令 2011/65/EU 的要求。

Diamond Diagnostics Inc. garante e declara que o(s) produto(s) listado(s) abaixo cumpre(m) o requisito do Regulamento de Dispositivos Médicos de Diagnóstico In Vitro EU 2017/46 e Diretiva 2011/65/EU.

Diamond Diagnostics Inc. гарантирует и заявляет, что продукты, перечисленные ниже, соответствуют требованиям Регламента ЕС 2017/46 о медицинских устройствах для диагностики in vitro и Директивы 2011/65/EU.

Vitro Diagnostica Medical Regulation 2017/746 and Directive 2011/65/EU المنتجات المذكورة أدناه تتوافق مع متطلبات الاتحاد الأوروبي المدرجة في
ن شركة دايموند دايانوسستكس تصرح و تؤكد أن التعليلة

Diamond Diagnostics Inc. garantisce e dichiara che i prodotti elencati di seguito sono conformi ai requisiti del Regolamento UE 2017/46 sui dispositivi medici per la diagnostica in vitro e della Direttiva 2011/65/EU.

Diamond Diagnostics, Inc. işbu belge ile aşağıda listelenen ürün(ler)in In Vitro Diagnostics Tıbbi Cihazlar Yönetmeliği EU 2017/746 ve Direktif 2011/65/EU gerekliliklerine uygun olduğunu garanti ve beyan eder.

Product(s) / Produkt(e) / Produit(s) / Producto(s) / 產品 (S) / Produto(s) / Продукт (ы) / المنتج (ق) / Prodott(i) /

Model: Abbott Architect Immunology, Abbott Cell Dyn Hematology

Electrodes & Accessories

Part Number:	Device Name:	Basic UDI-DI	Intended Use
AB-8921153401/1	Compressor Service Kit	081140301AB8921153401/1KS	Kit used to service the compressor within these instruments
AB-9216102D	Waste Sensor	081140301AB9216102DC9	Senses the level of the waste

Manufacturer's Name:

Diamond Diagnostics Inc.

Manufacturer's Address:

333 Fiske Street
Holliston, MA 01746
Tel: +1 (508) 429-0450

Diamond Diagnostics Inc. Hungary Branch Office
6 Oradna Street
1044 Budapest, Hungary
Tel: + 3617872222

(AR) Authorized Representative: Diamond Diagnostics Inc., Hungary Branch Office

(SRN) Single Registration Number: US-MF-000035435

Risk Class: Class A, Rule 5

Conformity Route: Annex IV, Self-Declared

Notified Body: N/A

Quality Systems Registration: ISO 9001:2015
ISO 13485:2016
EN ISO 9001:2015
EN ISO 13485:2016

Authorized Officer: Kathy Fisher **Date:** 6 May, 2024
Kathy Fisher
Director, Quality Assurance

