Collection and **Shipping Instructions**



aruplab.com/AAV5

500 Chipeta Way Salt Lake City, UT 84108-1221 Phone: 800-522-2787 Reference: DetectCDx

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AAV5 Total Antibody (TAb) Assay or AAV5 DetectCDx for ROCTAVIAN (Valoctocogene Roxaparvovec-Rvox) Eligibility in Hemophilia A

For In Vitro Diagnostic Use | Rx Only



The AAV5 DetectCDx™ assay is intended for the detection of anti-AAV5 antibodies in human plasma collected in 3.2% sodium citrate from adult patients with hemophilia A for whom ROCTAVIAN (valoctocogene roxaparvovec-rvox) treatment is being considered.

Ordering Instructions

The AAV5 DetectCDx test may be ordered electronically using a previously established laboratory information system (LIS) interface or ARUP Connect™, or the test may be ordered manually with a test requisition form.

- AAV5 DetectCDx requisition forms can be obtained through ARUP Client Services. Call 800-522-2787 for more information.
- If using a test requisition form, ensure all patient information is accurate and properly documented.

A shipping kit (ARUP #55016) containing the materials necessary to prepare plasma for shipment to ARUP Laboratories will be provided upon request. Visit aruplab.com/AAV5 or contact ARUP Client Services (800-522-2787) for more information.

Collection Instructions

- Collect the patient's whole blood in a 3.2% sodium citrate tube.
 - Samples that exceed 7.3% sodium citrate cannot be evaluated and may require patient redraw.
 - NOTE: When drawing blood for the AAV5 DetectCDx test, universal precautions for bloodborne pathogens should be observed.
- Centrifuge the specimen and separate the plasma within 72 hours of collection. Refer to your manufacturer's manual for recommended centrifuge speed and duration.
- Transfer 1 mL (minimum: 0.5 mL) of plasma into a pour-off (transport) tube. Failure to provide sufficient volume may result in the need for patient redraw.

- Label the transport tube with the patient's first and last name, date of birth, and sex.
- Freeze plasma specimen at -10°C or below.
- · Ship frozen plasma specimens to ARUP as soon as possible.
- NOTE: Plasma specimens must be completely frozen prior to shipment to ARUP Laboratories.

Shipping Instructions

All AAV5 DetectCDx specimens must be shipped to ARUP Laboratories in Salt Lake City, Utah.

- Print the appropriate paperwork (LIS interface packing list, ARUP Connect packing list, or manual requisition) and place in a specimen transport bag with the transport tube containing the frozen plasma.
- Place the frozen specimen transport bag in an insulated box containing dry ice.
- Ship the insulated box to ARUP Laboratories via your established courier service. Use overnight delivery to ensure next-day arrival at ARUP.
 - If you do not have an established courier service, use the FedEx waybill provided in the shipping kit.
 - If shipping via FedEx, affix the completed FedEx waybill to the outside of the insulated box.
- Arrange specimen pickup with your established courier or FedEx.

For additional instructions, please refer to the Instructions for Use. Visit **aruplab.com/AAV5** to learn more about AAV5 DetectCDx.

The AAV5 Total Antibody Assay for ROCTAVIAN (valoctocogene roxaparvovec-rvox) Eligibility in Hemophilia A ("AAV5 TAb Assay") or AAV5 DetectCDx is a qualitative in vitro diagnostic test by electrochemiluminescence intended for detection of antibodies in human plasma collected in 3.2% sodium citrate that bind to the adeno-associated virus servicy 6 [AAV5]. The AAV5 Tab Assay is indicated as an aid in the selection of abult hemophilia A patients for whom ROCTAVIAN treatment is being considered. Patients that are anti-AAV5 antibody positive (result of Detected) are not eligible for treatment with ROCTAVIAN, patients that are anti-AAV5 antibody negative (result of Not Detected) are eligible for treatment with ROCTAVIAN. This assay is for professional use and is a single-site assay performed at ARUP Laboratories.

Hemophilia A

AAV5 DetectCDx™

Visit **aruplab.com/AAV5** or contact your local account executive to learn more about AAV5 DetectCDx and ARUP Laboratories.



aruplab.com/AAV5

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Reference: DetectCDx

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AAV5 DetectCDx™ Explained

AAV5 DetectCDx is a prescription-only companion diagnostic (CDx) test that helps identify if patients with hemophilia A are eligible for the gene therapy treatment ROCTAVIAN (valoctocogene roxaparvovec-rvox).

ARUP is the only lab that offers the FDA-approved CDx assay for the gene therapy treatment ROCTAVIAN. We are a CAP-, ISO 15189-, and CLIA-certified reference lab with more than 35 years of experience supporting laboratories, physicians, and patients with unparalleled quality and service.

AAV5 DetectCDx is sponsored by BioMarin and offered at no cost to evaluate eligibility for an FDA-approved use. No patient, private health plan, government health program, or any other individual or entity shall be billed for this serotype test.⁶

This information is intended for laboratory us

How to Order

If you are a hospital, health system, or reference laboratory that is partnered with ARUP, testing will be available through ARUP Connect™ or can be ordered with a manual client requisition. Test orders and results may also be built to electronically interface with your medical record and laboratory information systems.

Otherwise, contact ARUP's Client Services at 1-800-522-2787 and reference DetectCDx to request the specimen shipping kit (**ARUP Supply Number 55016**).

Test Information

AAV5 Total Antibody (TAb) Assay or AAV5 DetectCDx for ROCTAVIAN (Valoctocogene Roxaparvovec-Rvox) Eligibility in Hemophilia A (3000959

Shipping

Ship the sealed kit via FedEx overnight or with an ARUP contracted courier. Visit **aruplab.com/AAV5** for more detailed collection and shipping instructions and to learn more about AAV5 DetectCDx.

^{1.} The AAVS Total Antibody Assay for ROCTAVIAN (valoctocogene roxaparvovec-rvox) Eligibility in Hemophilia A (YAAVS TAb Assay) or AAVS DetectCD is a qualitative in vitro diagnostic test by electrochemiluminescence intended for detection of antibodies in human plasma collected in 3.2% sodium citrate that bind to the adeno-associated virus serotype 5 (AAVS). The AAVS TAb Assay is indicated as an aid in the selection of adult hemophilia A patients for whom RIOCTAVIAN treatment is being considered. Patients that are anti-AAVS antibody positive (result of Detected) are not eligible for treatment with RIOCTAVIAN, patients that are anti-AAVS antibody negative (result of Not Detected) are eligible for treatment with RIOCTAVIAN, and its assay is for professional use and is a single-site assay performed at ARUP Laboratories. 2. For more detailed information about RIOCTAVIAN and its safety and efficacy, please on to biomarin-reaconnections could

^{3.} When drawing blood for the AAVS DetectCDX** Assay, universal presautions for bloodborne pathogens should be observed.
4. The stability of the patient sample during whole blood collection has been established for 72 h at room temperature or refrigerated (2 to 8 oC) in the AAV5 Tab Assay or AAV5 DetectCDX. Stability of the patient sample during processing has been established for 72 h at room temperature and 72 h refrigerated 2 to 8 oC) in the AAV5 DetectCDX. Patient samples with rheumatoid factor levels greater than 476 III/mL will interfere with the ability of the AAV5 DetectCDX to accurately detect anti-AAV5 antibodies. Patient samples with hylogendie levels greater than 500 mg/dL may interfere with the ability of AAV5 DetectCDX to accurately detect anti-AAV5 antibodies. Patient samples with hemoglobin levels greater than 800 mg/dL may interfere with the ability of AAV5 DetectCDX to accurately detect anti-AAV5 antibodies. Collected patient samples must not exceed 7.3% sodium cirtare. Patients with a Not Detected result may receive ROCTAVIAN infusion. 5. This test can only be conducted on plasma from adult male patients collected in 3.2% sodium cirtare anticoagulant. Other specimen types and conditions have not been validated. 6. Enrollment and eligibility requirements may apply and BioMarin reserves the right to terminate or amend this program without notice at any time.



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AAV5 DetectCDx

Step 2: Complete the test requisition form.

Step 1: Call ARUP Client Services (1 800 522 2787) to request a test requisition form.

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Date of Issuance: XXXXXXX





AAV5 Total Antibody Assay for ROCTAVIAN (valoctocogene roxaparvovec-rvox) Eligibility in Hemophilia A

Proprietary and established product name:

Device Trade Name: AAV5 DetectCDxTM

Device Generic Name: AAV5 Total Antibody (TAb) Assay for ROCTAVIAN (valoctocogene roxaparvovec-rvox) Eligibility in Hemophilia A

Intended use:

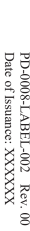
aid in the selection of adult hemophilia A patients for whom ROCTAVIAN treatment is being considered. citrate that bind to the adeno-associated virus serotype 5 (AAV5). The AAV5 TAb Assay is indicated as an electrochemiluminescence intended for detection of antibodies in human plasma collected in 3.2% sodium ROCTAVIAN; patients that are anti-AAV5 antibody negative (result of Not Detected) are eligible for Patients that are anti-AAV5 antibody positive (result of Detected) are not eligible for treatment with Hemophilia A ("AAV5 TAb Assay") or AAV5 DetectCDx is a qualitative in vitro diagnostic test by treatment with ROCTAVIAN. This assay is for professional use and is a single-site assay performed at The AAV5 Total Antibody Assay for ROCTAVIAN (valoctocogene roxaparvovec-rvox) Eligibility in ARUP Laboratories.

Contraindications:

None

Warnings and precautions:

- pathogens should be observed. When drawing blood for the AAV5 DetectCDxTM assay, universal precautions for bloodborne
- the AAV5 DetectCD x^{TM} to accurately detect anti-AAV5 antibodies. Patients with rheumatoid factor levels greater than 476 IU/mL will interfere with the ability for
- Patient samples with triglyceride levels greater than 500 mg/dL will interfere with the ability of the AAV5 DetectCDxTM to accurately detect anti-AAV5 antibodies





- the AAV5 DetectCD x^{TM} to accurately detect anti-AAV5 antibodies. Patient samples with Hemoglobin levels greater than 800 mg/dL will interfere with the ability of
- higher concentrations could not be evaluated. Patient samples collected for the AAV5 DetectCDxTM must not exceed 7.3% sodium citrate as
- is unknown. A positive assay result can occur due to detection of antibodies other than anti-AAV5 Cross-reactivity in the AAV5 DetectCDxTM assay to antibodies other than anti-AAV5 antibodies
- (typically indicative of a "Not Detected" result), that the sample still be considered "Detected." DetectCDx, it is recommended that if a sample with an SI value > 90 generates a CI value > 1.00 Since a potential prozone/hook effect was not evaluated for samples with SI > 90 with the AAV5

Summary and explanation of the test:

screening and confirmatory steps to reliably detect antibodies specific for AAV5 capsid. Patients evaluated are not eligible for treatment with ROCTAVIAN. for treatment with valoctocogene roxaparvovec-rvox (ROCTAVIAN) under the supervision of a physician. with the AAV5 DetectCDxTM who are anti-AAV5 antibody negative (result of Not Detected) are eligible citrated (3.2%) plasma specimens. The AAV5 DetectCDxTM uses a combination of concurrently conducted Patients evaluated with the AAV5 DetectCDxTM who are anti-AAV5 antibody positive (result of Detected) The AAV5 DetectCDxTM uses a bridging immunoassay to detect antibodies to AAV5 in human sodium

to insufficient quantities of FVIII or a dysfunctional FVIII. essential cofactor in the intrinsic coagulation cascade. This disorder can be inherited or acquired, leading 5,000 males. Hemophilia A is caused by deficiency in the activity of coagulation factor VIII (FVIII), an treatment for severe hemophilia A, an X-linked recessive bleeding disorder that affects approximately 1 in Valoctocogene roxaparvovec-rvox (ROCTAVIAN), or AAV5-hFVIII-SQ drug product, is a gene therapy

capsid mediates binding and uptake into cells, as well as trafficking to the cell nucleus. The vector genome plasma, synthesized from vector-transduced liver tissue. delivered by single intravenous dose and was designed to achieve stable expression of active FVIII in the transgene codes for an active form of FVIII that is used in the coagulation process. ROCTAVIAN is as ITRs). After unpackaging of the vector genome in the cell nucleus, recombination between the ITRs contains a transgene expression cassette inserted between the AAV DNA terminal sequences (referred to the SQ form of human FVIII (hFVIII-SQ) under the control of a liver-specific promoter. The AAV5 viral generates double-stranded, circular vector genomes that persist mainly as un-integrated episomes. The ROCTAVIAN is an adeno-associated virus serotype 5 (AAV5)-based gene therapy vector that expresses

demonstrated a titer-dependent reduction in transgene expression when dosed with AAV vectors (Scallan, can inhibit liver transduction and expression of the transgene product (Jiang, 2006, Blood); (Wang, 2011, detectable anti-AAV5 antibodies as determined by the AAV5 DetectCDx will be eligible for treatment with Since pre-existing anti-AAV5 antibodies may neutralize ROCTAVIAN, only patients who demonstrate no Hum Gene Ther), while immune-deficient mice reconstituted with purified human immunoglobulins ROCTAVIAN. The presence of neutralizing activity against AAV capsids in non-human primates (NHPs)



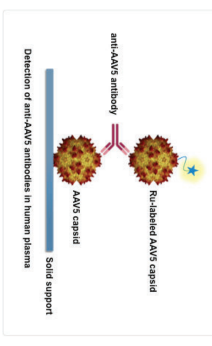
vectored Factor IX (FIX) transgene (Manno, 2006, Nature Med). suggested in the clinical setting by treatment of small numbers of hemophilia B patients with an AAV2-2006, Blood). Diminished efficacy, correlating with the presence of pre-existing immunity, has also been

step. The cut points for the screening and confirmatory assays were determined based on the statistical electrochemiluminescence (ECL) signal is specific. In the confirmatory step, samples are pre-incubated screening step and 1% false positive rate for the confirmatory step. analysis of a set of samples negative for anti-AAV5 antibodies yielding a 5% false positive rate for the antibodies that are present. If AAV5-binding antibodies are present, they will be bound by the unlabeled with unlabeled capsid (referred to as AAV5 Confirmatory Reagent) to compete for any anti-AAV5 assesses for the presence of anti-AAV5 antibodies, while the confirmatory step determines if the screening and confirmatory steps to reliably detect antibodies specific for AAV5 capsid. The screening step citrated (3.2%) plasma specimens. The AAV5 DetectCDxTM uses a combination of concurrently conducted AAV5 capsid, resulting in a reduced ECL signal for the confirmatory step as compared to the screening The AAV5 DetectCDxTM uses a bridging immunoassay to detect antibodies to AAV5 in human sodium

Principles of the test procedure:

500 Chipeta Way, Salt Lake City, UT 84108 The AAV5 DetectCDxTM is to be performed only at ARUP Laboratories, a single laboratory site located at

specimen, the plate is washed, and SULFO-TAG AAV5-CMV-GFP capsid is added to each well. Antiadded in duplicate to specific wells of the plate. If anti-AAV5 antibodies are present in the specimen, they substrate is added to each well. The plate is read on a research use only (RUO) ECL-based plate reader. in an electrochemiluminescence (ECL) reaction. After incubation and washing, tripropylamine (TPA) will bind to the unlabeled AAV5-CMV-GFP capsid coating the wells. After incubation with patient with assay diluent containing casein, and washed again. The patient plasma specimen is diluted and then A MULTI-ARRAY® 96-well plate is coated with unlabeled AAV5-CMV-GFP capsid, washed, blocked Each well of the plate is electrically stimulated and the resultant ECL signal is measured. AAV5 antibodies bind to SULFO-TAG capsid (also referred to as ruthenylated capsid), which participates



ruthenylated AAV5 capsid. The Ru-label participates in the generation of an electrochemiluminescent signal Figure 1. Anti-AAV5 antibody forms a bridge between AAV5 capsid coating the immunoassay plate and that indicates the presence of anti-AAV5 antibodies.



LPC must meet the pre-established criteria for the between-well coefficient of variation (CV) for replicate control (LPC), and a high antibody positive control (HPC). For run acceptance, the NEG, CC, HPC, and within the established acceptance range. wells. The HPC and LPC must screen and confirm positive, and the HPC, LPC, and NEG signals must fall Each 96-well plate includes a cut point control (CC), negative control (NEG), a low antibody positive

obtained for the SI and CI (see Figure 2): considered if anti-AAV5 antibodies are not detected in the screening step. Results are based on the values for the confirmatory and screening assays and dividing this by the confirmatory cut point. The CI is not Confirm Index (CI). The confirm index (CI) is obtained by calculating the ratio of mean signals obtained normalized screening result by the screening cut point. Results for the confirmatory step are expressed as a Results for the screening step are expressed as a Screen Index (SI). The SI is calculated by dividing the

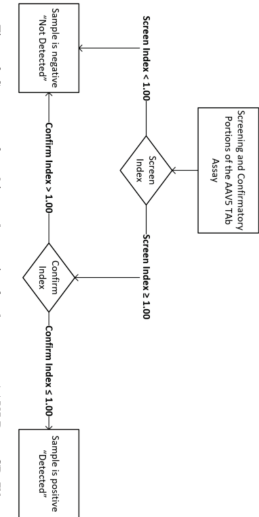


Figure 2. Summary of resulting and reporting for the two-step AAV5 DetectCDxTM

antibodies. Specimens with SI < 1.00, or SI \geq 1.00 with a CI > 1.00, are reported as Not Detected for anti-AAV5

Specimens with $SI \ge 1.00$ and $CI \le 1.00$ are reported as Detected for anti-AAV5 antibodies

Reported Results:

Patients evaluated with the AAV5 DetectCDxTM who are anti-AAV5 antibody negative (result of Not will be sent to the designated physician. The following are the standard report results: Upon completion of testing at ARUP Laboratories, a test report with the results of the AAV5 DetectCDxTM Detected) are eligible for treatment with ROCTAVIAN under the supervision of a physician



- rvox) Detected: patient is not eligible for treatment with ROCTAVIAN (valoctocogene roxaparvovec
- rvox) Not Detected: patient is eligible for treatment with ROCTAVIAN (valoctocogene roxaparvovec

Reagents:

The primary reagents for the AAV5 DetectCDxTM include:

AA	AAV5 DetectCDxTM Reagents and Storage Conditions	nd Storage Conditions
Reagent	Storage Conditions	Component(s)
AAV5 Coated Plate Set	Refrigerated (2 to 8 °C)	 MULTI-ARRAY® 96-well plate
		 AAV5 Confirmatory Reagent
		AAV5 Detection Reagent
AAV5 Control Set	Frozen (-70 °C)	Low positive control
		High positive control
		• Cut point control
		Negative control
Read Buffer T (1X)	Room temperature (20 to Read Buffer T (1X)	Read Buffer T (1X)
	25 °C)	

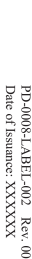
Additional reagents used in the AAV5 DetectCDx:

- TBS Buffer (1X) with 1% casein
- 1X DPBS
- Tween 20 (proteomics grade), 1.0% (v/v)
- ProClin 300, 0.05% (v/v)

QuickPlex SQ 120 instrument, as identified by a specific serial number. **Instruments:** The MULTI-ARRAY® 96-well plate used in the AAV5 DetectCDxTM is read on a MESO

General laboratory instruments and materials that are also used in the AAV5 DetectCDxTM include:

- Refrigerator capable of 2 to 8 °C
- Freezers capable for -10 °C or colder and -70 °C or colder
- Single-channel pipette set
- Multi-channel pipettes
- Vortex mixer
- Minicentrifuge capable of 400 RPM
- Microplate shaker
- Microplate washer
- Microplate adhesive film
- PCR aluminum sealing film
- 0.2 mL and 1.2 mL 8-well strip tubes with cap





Workbench® version 4.0 and Cerner's Millennium Helix Unified Case Manager® version 2018.13.02 Software: The software used with the AAV5 DetectCDxTM is comprised of the MSD Discovery

Specimen shipping kit contents:

specimen shipping kit includes the following components: physician. Use of the shipping kit is optional; the kit will be provided to customers upon request. The A specimen shipping kit provided by ARUP Laboratories may be used by the ordering laboratory or

- Tube, Round False Bottom, Std Transport, 4mL screw cap
- Collection and Shipping Instructions
- Bag, ARUP Sample, Frozen, Ziploc w/ Absorbent pad
- Box, Shipping (cardboard with foam cooler insert; comes with a UN3373 and dry ice label)
- FedEx Priority Pre-Paid Shipping Label

Specimen collection and shipping:

in specimen rejection and will not be tested. must be transported to ARUP Laboratories frozen on dry ice. Any specimens not received frozen will result ordering interface (available only to existing ARUP clients). The TRF must be fully completed. All samples To order the AAV5 DetectCDx[™] assay, use the ARUP test requisition form (TRF) or ARUP's web-based

Collection Instructions

- Collect the patient's whole blood in a 3.2% sodium citrate tube.
- Samples that exceed 7.3% sodium citrate cannot be evaluated and may require patient redraw.
- pathogens should be observed. NOTE: When drawing blood for the AAV5 DetectCDx test, universal precautions for bloodborne
- manufacturer's manual for recommended centrifuge speed and duration. Centrifuge the specimen and separate the plasma within 72 hours of collection. Refer to your
- ARUP Transport Tube (polypropylene). Sample stability for the AAV5 DetectCDxTM has not been evaluated in tube types other than the Transfer 1 mL (minimum: 0.5 mL) of plasma into a polypropylene pour-off (transport) tube.
- Failure to provide sufficient volume may result in the need for patient redraw
- Label the transport tube with the patient's first and last name, date of birth, and sex
- Freeze plasma specimen at -10°C or below.
- shipped to ARUP Laboratories. to ensure next day arrival at ARUP. NOTE: Plasma specimens must be frozen before they are Ship frozen plasma specimens to ARUP as soon as possible on dry ice and use overnight delivery
- Plasma samples can be stored frozen (-10 to -70°C) for up to 12 months. Minimize number of freeze/thaw events, not to exceed 6 events.



collection, and shipping samples to ARUP Laboratories. shipping kit or found online at http://www.aruplab/aav5 for further details about test ordering, specimen Please refer to the AAV5 DetectCDxTM Collection and Shipping Instructions in the optional specimen

Limitations:

laboratory site located at 500 Chipeta Way, Salt Lake City, UT, 84108 This assay is intended for professional use only and is to be performed only at ARUP Laboratories, a single

- For in vitro diagnostic use
- For professional use only
- For prescription use only
- This test is intended to be performed on specific serial number-controlled instruments at ARUP Laboratories

Performance characteristics:

Detection capabilities:

qualification procedures. The detection capability of the AAV5 DetectCDxTM has been defined for internal quality control and

Precision:

Measurement Procedures and CLSI EP12-A2 - User Protocol for Evaluation of Qualitative reagents. The precision studies were based on CLSI EP05-A3 - Evaluation of Precision of Quantitative below Performance. AAV5 DetectCDx precision was assessed using five sample types, as indicated in the table The precision of the AAV5 DetectCDxTM assay was evaluated across days, operators, instruments and Test

	Sample Typ	Sample Types Used in DetectUDx ^{1,M} Precision Evaluation	Precision Eval	uation
Sample Type		SI Value		CI Value
	Target	Measured (mean)	Target	Measured (mean)
High negative	< 1.00	0.87	~ 1.20	1.193
Cutoff	> 1.00	1.04	~ 1.00	1.005
Low positive	> 1.00	1.56	$\sim \! 0.80$	0.695
Mid positive	~1.80	1.95	~0.60	0.538
High positive	> 10.0	40.01	< 0.20	0.0360

variations do not impact assay results. DetectCDxTM is acceptable and that operator-to-operator, instrument-to-instrument, and reagent lot-to-lot Results (summarized in the table below) indicate that inter- and intra-assay precision in the AAV5



Experimental	Runs	Qualitative Agreement	% Coefficient
Conditions			of Variance
Single operator,	2 runs per day	100% QA for high	$\%CV \le 15\%$ for
single instrument,	20 test days	negative, low positive,	all sample types
single raw material	2 replicates per	mid positive and high	tested
reagent lot	sample	positive samples	
Single operator,		100% QA for high	$\%CV \le 15\%$ for
single instrument,	16 replicates per	negative, low positive,	all sample types
single raw material	sample	mid positive and high	tested
reagent lot		positive samples	
3 operators, 1	1 run per day per	100% QA for high	$\%CV \le 15\%$ for
instrument, 1	operator	negative, low positive,	all sample types
production reagent	5 test days	mid positive and high	tested
lot	5 replicates per sample	positive samples	
1 operator, 2	1 run/day	100% QA for high	$\%CV \le 15\%$ for
instruments, 1	5 test days	negative, low positive,	all sample types
production reagent	5 replicates per	mid positive and high	tested
lot	sample	positive samples	
1 operator, 1	1 run/day	100% QA for high	Between-lot
instrument, 3	6 test days	negative, low positive,	%CV < 15%
vendor reagent lots	4 replicates per	mid positive and high	
	sample	positive samples	
	Conditions Conditions Conditions Conditions Single operator, single instrument, single raw material reagent lot Single operator, single raw material reagent lot 3 operators, 1 instrument, 1 production reagent lot 1 operator, 2 instruments, 1 production reagent lot 1 operator, 1 instrument, 1 production reagent lot 1 operator, 2 instruments, 1 production reagent lot 1 operator, 1 instrument, 3 vendor reagent lots	Conditions Conditions Conditions Conditions Conditions 2 runs per day 20 test days 2 replicates per days 2 replicates per day 3 operators, 1 dereplicates per day 3 operator day 4 replicates per day 5 test days 5 replicates per day 5 replicates per day 5 replicates per day 5 replicates per day 6 test days 5 replicates per day 6 test days 7 replicates per day 8 per day 9 pe	al Runs 2 runs per day 1, 20 test days rial 2 replicates per sample 1 fo replicates per rial sample 1 run per day per operator 5 test days 5 replicates per sample 1 run/day 5 test days 5 replicates per sample 1 run/day 6 test days 6 test days 6 test days 4 replicates per sample 1 run/day 6 test days

analysis is presented in the table below. for the cutoff sample from the precision studies. The observed results (n=308) were determined as Detected the performance of the assay at sample near cutoff was as expected. A total of 308 results were generated CI: (37%; 48%). The sample near cutoff performed as expected. Details of AAV5 DetectCDx sample 132 times and determined as Not Detected 176 times. The percent Detected was 43% (132/308) with 95% The data for the sample near cutoff (SI > 1.00, CI \sim 1.00) was collected and evaluated to determine whether

Results of Sample Near AAV5 DetectCDx TM	e Near AAV5 De	tectCD _X TM
	SI	CI
n	308	308
Mean	1.04	1.01
Median	1.04	1.00
SD	0.054	0.060
%CV	5.2%	6.0%

Interference:

plasma) and exogenous substances (e.g. common over-the-counter medicines, prescription drugs). interference study evaluated the impact of substances on the assay results using three sample types that Interference testing was based on CLSI EP07-A3 - Interference Testing in Clinical Chemistry, 3rd Edition; CLSI EP37-ED1 - Supplemental Tables for Interference Testing in Clinical Chemistry, 1st Edition. The AAV5 DetectCDxTM was evaluated for interference by endogenous (naturally present in human



corresponded to a high negative sample, a low positive sample, and a high positive sample, as indicated in the tables below.

Sam	ple Types Used	Sample Types Used in Endogenous & Exogenous Substances Evaluation	nous Substanc	es Evaluation
Sample Type		SI Value		CI Value
	Target	Measured (mean)	Target	Measured (mean)
High negative	< 1.00	0.850	~ 1.20	1.278
Low positive	> 1.00	1.260	~ 0.80	0.862
High positive	> 10.0	23.670	< 0.20	0.048

Rheumatoid factor (RF) interference was tested by evaluating the change in AAV5 DetectCDx assay results of rheumatoid factor. when a low positive sample was added to a high negative sample in the presence of different concentrations

changed the qualitative output of the sample compared to control. A substance was also considered an and below the critical assay cutoff, compared to control were > 10% with a high degree of confidence. interferent if the change in the SI/CI values of the high negative or low positive sample, samples above A substance was considered an interferent to the AAV5 DetectCDxTM if addition of the test substance

	Interfering Substances to AAV5 DetectCDx	ectCDx
Substance	Test concentration	Impact on
		Qualitative Test Result
Hemoglobin	1000 mg/dL	Could convert sample to Not
		Detected result
Triglycerides	750 mg/dL	Could convert sample to Not
		Detected result
Rheumatoid Factor	1285 IU/mL, 1750 IU/mL, 3695 IU/mL No expected impact	No expected impact

compared to control † RF interfered with the AAV5 DetectCDx in a dose-dependent manner with > 10% difference in assay values

Non-interfering Endogenous and Exogenous Substances*	Exogenous Substances*
Substance	Test concentration
Albumin	6 mg/dL
Bilirubin, conjugated	40 mg/dL
Bilirubin, unconjugated	40 mg/dL
Triglycerides	$500 \mathrm{mg/dL}$
Triglycerides	$200 \mathrm{mg/dL}$
Hemoglobin	$800 \mathrm{mg/dL}$
Hemoglobin	$400 \mathrm{mg/dL}$
Rheumatoid Factor	476 IU/mL
Acetaminophen	15.6 mg/dL
Advate	384 IU/dL
Atazanavir	1.95 mg/dL
Atorvastatin	$0.075 \mathrm{mg/dL}$
Bictegravir	1.85 mg/dL
Biotin	0.351 mg/dL
Doravirine	$0.289 \mathrm{mg/dL}$



Non-interfering Endogenous and Exogenous Substances*	Exogenous Substances*
Substance	Test concentration
Eloctate	324 IU/dL
Fexofenadine	$0.116 \mathrm{mg/dL}$
Hemlibra	170 μg/mL
Hemofil-M	150 IU/dL
Heparin	330 IU/dL
Ibuprofen	21.9 mg/dL
Lisinopril	0.0246 mg/dL
Naproxen	36.0 mg/dL
Omeprazole	0.84 mg/dL
Oxycodone	0.0324 mg/dL
Sodium citrate†	7.3% (short draw)
Tenofovir	0.0978 mg/dL
Vitamin C	5.25 mg/dL

Higher concentrations of sodium citrate could not be evaluated with the AAV5 Detect CDx^{TM}

Cross-reactivity with other antibodies:

unknown. A positive assay result can occur due to detection of antibodies other than anti-AAV5 antibodies. Cross-reactivity in the AAV5 DetectCDxTM assay to antibodies other than anti-AAV5 antibodies is

Prozone effect:

sample for eight (8) dilution steps to cover the range from high positive to negative Screen Index and created by diluting the high titer positive AAV5 plasma samples into the anti-AAV5 negative plasma identified in historical studies conducted at ARUP Laboratories. Individual two-fold dilution series were antibody produce a prozone (hook) effect. The study samples utilized distinct plasma samples from three The AAV5 DetectCDxTM was evaluated to determine whether elevated concentrations of anti-AAV5 Confirm Index values. (3) non-hemophilia A donors that represent the highest AAV5 titer positive samples that were previously

of ~90. Human specimens with SI values greater than 90 were not evaluated in this study and may exhibit a prozone effect. The results from this test indicate that a prozone effect was not observed for samples with starting SI values

Carryover:

The possibility of carryover and well-to-well cross-talk was evaluated for the AAV5 DetectCDxTM assay. high positive samples. The study sample set indicated in the table below was used to create an alternating pattern of negative and

the effect of these substances on the assay is unknown * Cholesterol and Celebrex have not been evaluated as potential interferents to the AAV5 DetectCDxTM assay, so



Sample Type	npie Types Ex	SI Value SI Value CI Value	Id Cross-1 alk n	CI Value
Sampic Type		OT A UTUC		CIVAIUC
J	Target	Measured (mean)	Target	Measured (mean)
Negative	< 1.00	0.88	>1.00	1.427
High positive	50-85	49.40	0.03 - 0.15	0.026

sections of the plate. assay modes and the negative and high positive samples were swapped between plates to address all The two (2) AAV Coated Plates were arranged so that the locations of the screening and confirmatory

100% Detected	High positive
100% Not Detected	Negative
Reported values	Sample Type
Summary of AAV5 DetectCDx TM Carryover and Cross-Talk Evaluation	Summary of AAV5 Detect(

Based on these results, it was concluded that carryover or well-to-well cross-talk were not observed in the

Stability:

impact of various storage and transport conditions of reagents and human whole blood and plasma using three sample types that corresponded to a high negative sample, a low positive sample, and a high positive on CLSI EP25-A - Evaluation of Stability of In Vitro Diagnostic Reagents. Stability studies evaluated the Stability of the reagents, collections, and samples for the AAV5 DetectCDxTM assay were evaluated based sample, as indicated in the table below.

	San	Sample Types Evaluated in Stability Tests	tability Tests	
Sample Type		SI Value		CI Value
	Target	Measured (mean)	Target	Measured (mean)
High negative	< 1.00	0.89	~ 1.20	1.245
Low positive	> 1.00	1.46	$\sim \! 0.80$	0.768
High positive	> 10.0	31.08	< 0.20	0.038

degree of confidence. sample, samples above and below the critical assay cutoff, compared to control were > 10% with a high the results of the AAV5 DetectCD x^{TM} if the change in the SI/CI values of the high negative or low positive the sample compared to control was changed. A condition and/or timepoint was also considered to impact A condition and/or timepoint was considered to impact the AAV5 DetectCDxTM if the qualitative output of

AAV5 DetectCDx TM F	AAV5 DetectCDx TM Plasma Sample Stability
Storage Conditions	Stability
Room temperature (20 to 25 °C)	72 hours
Refrigerated (2 to 8 °C)	28 days
Frozen (-10 °C or colder)	12 months
Frozen (-70 °C or colder)	12 months
Freeze-thaw cycles	7 events



AAV5 DetectCDx TM Sa	AAV5 DetectCDx TM Sample Collection Stability
Conditions	Stability
Plasma, room temperature (20 to 25 °C)	72 hours
Plasma, refrigerated (2 to 8 °C; for storage post-	72 hours
processing prior to freezing)	
Whole blood, room temperature (20 to 25 °C)	72 hours
Whole blood, refrigerated (2 to 8 °C; for storage	72 hours
prior to processing to plasma)	

AAV5 DetectCDx TM Plasma	AAV5 DetectCDx TM Plasma Sample Transport Stability
Transport Conditions	Stability
Refrigerated (with gel packs)	10 days
Ambient temperature	10 days
Frozen (on dry ice)	10 days
Elevated temperature $(37 ^{\circ}\text{C} \pm 2 ^{\circ}\text{C})$	1 day
Frozen (on ice pack)	7 days

AAV5	AAV5 DetectCDxTM Established Reagent Stability	t Stability
Reagent(s)	Conditions	Stability
AAV5 Plate Components	Frozen (-70 °C)	12 months
AAV5 Run Control Set	Frozen (-20 °C)	12 months
AAV5 Coated Plate Set	Refrigerated (2 to 8 °C)	7 days
Read Buffer T (1X)	20 to 25 °C	12 months

Expected Values

Patient Population Demographics:

AAV5 DetectCDx assay results (Detected vs Not Detected). A number of patient population demographic variables were analyzed for their potential association with

Percent of Detected AAV5 DetectCDx Results Stratified by Race and Ethnicity	Results Stratifi	ed by Race and Ethnicity
Race	N	Percent Detected
White	618	27.8% (172/618)
Asian	159	28.3% (45/159)
Black or African American	110	34.5% (38/110)
Native Hawaiian or other Pacific Islander	2	0.0% (0/2)
Not Provided or Multiple	138	40.6% (56/138)
Combined	1,027	30.3% (311/1,027)
Ethnicity		
Hispanic or Latino	27	29.6% (8/27)
Not Hispanic or Latino	965	29.8% (288/965)
Not provided	35	42.9% (15/35)
Combined	1,027	30.3% (311/1,027)

Higher seropositivity (percent of results Detected) was observed for the "Black or African American" group (34.5% Detected).





Percent of Detected AAV5 De	etectCDx Results S	Percent of Detected AAV5 DetectCDx Results Stratified by Country of Origin
Country of Origin	N	Percent Detected
Australia	45	15.6% (7/45)
Belgium	19	21.1% (4/19)
Brazil	102	32.4% (33/102)
France	116	37.1% (43/116)
Germany	101	25.7% (26/101)
Israel	12	8.3% (1/12)
Italy	24	33.3% (8/24)
South Africa	112	35.7% (40/112)
Spain	14	21.4% (3/14)
South Korea	6	33.3% (2/6)
Taiwan	40	35.0% (14/40)
United Kingdom	94	18.1% (17/94)
United States	168	28.0% (47/168)
Russia	91	46.2% (42/91)
Japan	84	29.8% (25/84)
Combined	1,028	30.4% (312/1,028)

observed in Israel (8%) and United Kingdom (18.1%). A high level of seropositivity (percent results Detected) was observed in Russia (46%) and a low level was

Percent of Detected AAV5 DetectCDx Results Stratified by Type of FVIII Replacemen	Dx Results Stratil	fied by Type of FVIII Replacement
	N	Percent "Detected"
On demand	108	45.4% (49/108)
Prophylaxis	891	26.4% (235/891)
Combined	999	28.4% (284/999)

prophylaxis group. The "on-demand" group experienced a higher seropositivity rate (percent results Detected) than the

Summary of clinical study:

Study Design:

and effectiveness of the AAV5 DetectCDx for its intended use was demonstrated through testing of interventional studies. The AAV5 DetectCDx was utilized in five (5) of these clinical studies. The safety evaluated at ARUP Laboratories in Salt Lake City, Utah using the AAV5 DetectCDxTM assay. the safety and efficacy of ROCTAVIAN; ClinicalTrials.gov Identifier NCT03370913). Samples were all specimens from hemophilia A patients enrolled in the clinical study 270-301 (study objective to evaluate The ROCTAVIAN clinical development program consists of six (6) interventional studies and two (2) non-

Study Population Demographics

selection of hemophilia A patients for treatment with ROCTAVIAN is based on the 134 patients from plasma collected in 3.2% sodium citrate samples for the presence of AAV5 antibodies to aid in the The effectiveness of the AAV5 DetectCDx as a companion diagnostic device for the testing of human



replacement therapy. There were no subjects on Emicizumab prophylaxis. study 270-301, who had a "Not Detected" AAV5 DetectCDx result and were enrolled in the clinical except two (2) subjects were HIV negative. Subjects were previously treated only with prophylactic FVIII study. In study 270-301, 134 subjects, aged 18 to 70 years (median: 30 years), received ROCTAVIAN The population was 72% White (96 patients), 14% Asian (19 patients), and 11% Black (15 patients). All

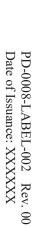
Demographics of 270-301 study population	01 study population
Age at enrollment, years	
Mean (SD)	31.7 (10.3)
Median (Range)	30.0 (18, 70)
Sex, n (%)	
Male	134 (100)
Race, n (%)	
Asian	19 (14.2)
Black or African American	15 (11.2)
Native Hawaiian or other Pacific Islander	1 (0.7)
White	96 (71.6)
Not provided due to patient privacy	3 (2.2)
Ethnicity, n (%)	
Hispanic or Latino	7 (5.2)
Not Hispanic or Latino	127 (94.8)
Type of FVIII treatment for hemophilia A, n (%)	
Prophylaxis	134 (100)

Study Results:

effectiveness of ROCTAVIAN. The results from this study support the clinical benefit of the AAV5 analysis met the pre-specified NI margin in the efficacy evaluable study subjects, indicating the compared with ABR during the baseline period with the NI margin set at 3.5 bleeds per year. The NI annualized bleeding rate (ABR) in the efficacy evaluation period following ROCTAVIAN administration dose of ROCTAVIAN. The primary efficacy outcome was a non-inferiority (NI) test of the difference in ROCTAVIAN. Adult hemophilia A patients in the study received a single administration of 6E13 vg/kg anti-AAV5 antibodies as an aid in the selection of hemophilia A patients for treatment with DetectCDx in the selection of hemophilia A patients for treatment with ROCTAVIAN. The results from study 270-301 support the clinical benefit of the AAV5 DetectCDx in the detection of

one (1) clinical study (270-301) on 134 subjects with severe hemophilia A exposed to ROCTAVIAN. selection of hemophilia A patients for treatment with ROCTAVIAN is based on the data generated in plasma collected in 3.2% sodium citrate samples for the presence of AAV5 antibodies to aid in the follow-up of 66 weeks and a median follow-up of 162 weeks (range 66 to 255 weeks). All subjects received a single dose of $6 \times 10^{13} \text{ vg/kg}$ of body weight of ROCTAVIAN with a minimum The safety evaluation of AAV5 DetectCDx as a companion diagnostic device for the testing of human

no participants discontinued from the study as a result of a treatment emergent adverse event (TEAE). The benefit-risk assessment. All subjects successfully completed their full-dose infusion of ROCTAVIAN, with ROCTAVIAN was found to have an acceptable safety and tolerability profile that supports a positive





upper limit of normal (ULN). Refer to the ROCTAVIAN Full Prescribing Information for more phosphokinase (CPK), factor VIII activity levels, gamma glutamyl transferase (GGT), and bilirubin above aminotransferase (ALT), aspartate aminotransferase (AST), lactate dehydrogenase (LDH), creatine most common adverse reactions to ROCTAVIAN were rash, headache, nausea, fatigue, diarrhea, and lab abnormalities. The most common laboratory abnormalities to ROCTAVIAN were

ongoing clinical trials and proposed post-approval studies Long-term safety of ROCTAVIAN continues to be monitored as outlined in the risk management plan in

Refer to Drugs@FDA for the most recent ROCTAVIAN product labeling

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