

# **Declaration of Conformity**

**Certificate Identification:** Legal Manufacturer's Name: Legal Manufacturer's Address: DoC-4P5220, 4P5201, 4P5211-SD DELK Abbott GmbH & Co. KG Max-Planck-Ring 2, 65205 Wiesbaden, Germany

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
4P52-20	59090	Hemoglobin A1c Reagent	Self-declared
4P52-01	53315	Hemoglobin A1c Calibrator	Self-declared
4P52-11	44435	Hemoglobin A1c Controls	Self-declared

Authorized European Representative (name and address)	N/A
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038, USA.
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:

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**Diana Romero** 

Position:

Full Name:

**Director**, Site QA

Date of Approval:

17-NOV-2017

Signature:

Full Name:

Position:

Mark Littlefield

Assoc. Director, Regulatory Affairs

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Date of Approval:

17-NOV-2017

Date Issued:

17-100-2017

65205 Wiesbaden, Germany

Place Issued:

Supersedes:

N/A

Effective (Date or Lot Number):

17-Nov-2017

# HEMOGLOBIN A10 CALIBRATORS

This package insert contains information for the use of Hemoglobin A1° Calibrators on the ARCHITECT  $c\,8000$  and  $c\,4000$  Systems.

### Read Highlighted Changes: Revised April 2017.

Package insert instructions must be carefully followed. Reliability of assay results cannot be guaranteed if there are any deviations from the instructions in this package insert.

### INTENDED USE

For use in the calibration of the Hemoglobin A<sub>1c</sub> assay on the ARCHITECT c 8000 and c 4000 Systems.

Refer to the Hemoglobin A1c reagent package insert and the ARCHITECT System Operations Manual for additional information.

### **CONTENTS / MATERIALS PROVIDED**

REF 4P52-01 Hemoglobin A1c Calibrators

CAL 1 1 x 1.6 mL

CAL 2 1 x 1.6 mL

Hemoglobin A<sub>1°</sub> Calibrators (lyophilized) contain hemoglobin and glycated hemoglobin from human whole blood. Prior to lyophilization, the calibrator matrix is an MES-buffered solution. **CAL** 1 and **CAL** 2 contain ofloxacin as a preservative.

# The value-assigned $A_{1\circ}$ Calibrator values are within the following hemoglobin $A_{1\circ}$ ranges:

	CAL 1	CAL 2
Hemoglobin A1c Range	4.59 to 6.02 %HbA <sub>1c</sub>	10.52 to 13.37 %HbA1c

Actual analyte concentrations for this lot of calibrators are listed in the Hemoglobin  $A_{1c}$  Calibrator value sheet, packaged with the calibrators. Verify that the lot number listed on each calibrator bottle agrees with the lot number printed on the value sheet.

### STANDARDIZATION

Each lot of calibrators is value-assigned. The concentration of glycated hemoglobin (HbA<sub>1c</sub>) and total hemoglobin (THb) is provided for each lot. Calibrators are prepared gravimetrically, lyophilized, and then value assigned using secondary calibrators that are traceable to the International Federation of Clinical Chemistry and Laboratory Medicine (IFCC) reference method.

### MATERIALS REQUIRED BUT NOT PROVIDED

- Calibrated pipette capable of measuring 1.6 mL
- Purified or deionized water
- · Vortex (optional)

### PRECAUTIONS

### **Precautions for Users**

- · IVD
- For In Vitro Diagnostic Use.
- · Rx ONLY
- · Do not use components beyond the expiration date.
- Do not mix components from different kit lot numbers.



CAUTION: This product contains human-sourced and/or potentially infectious components. Refer to the CONTENTS / MATERIALS PROVIDED section of this package insert. No known test method can offer complete assurance that products derived from human sources or inactivated microorganisms will not transmit infection. Therefore, all human-sourced materials should be considered potentially infectious. It is recommended that these calibrators and human specimens be handled in accordance with the OSHA Standard on Bloodborne Pathogens.<sup>1</sup> Biosafety Level 2<sup>2</sup> or other appropriate biosafety practices<sup>3,4</sup> should be used for materials that contain or are suspected of containing infectious agents. The human blood used in the calibrators is nonreactive for HBsAg, HIV-1 RNA or HIV-1 Ag, anti-HCV, and anti-HIV-1/ HIV-2.

A1c Calibrators REF 4P52-01 307261/R03 S4P5X0 FOR USE WITH ARCHITECT

The following warnings and precautions apply to: CAL 1 and CAL 2

	WARNING:	
>	H319	Causes serious eye irritation.
	H402*	Harmful to aquatic life.
	Prevention	
	P264	Wash hands thoroughly after handling.
	P273*	Avoid release to the environment.
	P280	Wear protective gloves / protective clothing / eye protection.
	Response	
	P305+P351+ P338	IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
	P337+P313	If eye irritation persists: Get medical advice / attention.
	Disposal	
	P501*	Dispose of contents / container in accordance with local regulations.

\* Not applicable where regulation EU 1272/2008 (CLP) has been implemented.

- Safety Data Sheets are available at www.abbottdiagnostics.com or contact your local representative.
- For a detailed discussion of safety precautions during system operation, refer to the **ARCHITECT System Operations Manual**, *Section 8.*

### STORAGE AND STABILITY

Unopened, lyophilized Hemoglobin A<sub>1c</sub> **CAL1** and **CAL2** are stable until the expiration date when stored at 2 to 8°C.

Store reconstituted calibrators at 2 to 8°C and use within 10 days. Do not freeze.

### PREPARATION OF CALIBRATORS

- The lyophilized calibrators should be reconstituted as follows:
- 1. Add 1.6 mL of purified or deionized water.
- Close bottle and allow the contents of the bottle to sit at room temperature for at least 10 minutes.
- 3. Mix thoroughly by low speed vortexing or by gently inverting 10 times before use. Avoid the formation of foam.

NOTE: After reconstitution, the calibrators do not require pretreatment.

### INSTRUCTIONS FOR USE

To perform a calibration:

- Enter the calibrator values for HbA1c and THb into the calibration file from the Hemoglobin A1c Calibrator value sheet.
   IMPORTANT: Calibrator values are defined in the Configure calibrator set screen. For the Hemoglobin A1c assay, it is also necessary to configure the Blank concentration in the Configure assay parameters - Calibration screen.
- Mix calibrators several times by gently inverting to ensure homogeneity. Avoid the formation of foam.
- 3. Open the bottles, place an appropriate amount of each calibrator into a separate sample cup.
- 4. Cap bottles and store at 2 to 8°C after use.
- Remove air bubbles, if present in the sample cup, with a new applicator stick. To minimize volume depletion, do not use a transfer pipette to remove the bubbles.

**CAUTION:** Bubbles in calibrators may interfere with proper detection of calibrator level in the sample cup causing insufficient calibrator aspiration that could impact results.

- 6. Place each sample cup in the assigned position.
- 7. Perform calibration as indicated in *Section 6* of the **ARCHITECT** System Operations Manual.

### INDICATIONS OF INSTABILITY OR DETERIORATION

Instability or deterioration should be suspected if there are visible signs of leakage, extreme turbidity, microbial growth, or if calibration does not meet the appropriate reagent package insert and/or **ARCHITECT System Operations Manual** criteria.

### LIMITATIONS OF THE PROCEDURE

This assay must be performed by qualified laboratory personnel, under appropriate laboratory conditions, solely for the intended use of the assay.

### **BIBLIOGRAPHY**

- 1. US Department of Labor, Occupational Safety and Health Administration. 29 CFR Part 1910.1030. *Bloodborne Pathogens*.
- US Department of Health and Human Services. *Biosafety in* Microbiological and Biomedical Laboratories, 5th ed. Washington, DC: US Government Printing Office, December 2009.
- 3. World Health Organization. *Laboratory Biosafety Manual*, 3rd ed. Geneva: World Health Organization, 2004.
- Clinical and Laboratory Standards Institute (CLSI). Protection of Laboratory Workers From Occupationally Acquired Infections; Approved Guideline—Fourth Edition. CLSI Document M29-A4. Wayne, PA: CLSI; 2014.

### TRADEMARKS

ARCHITECT, c8000, and c4000 are trademarks of Abbott Laboratories in various jurisdictions.

All other trademarks are property of their respective owners.

Key to Symbols		
CAL	Calibrator	
CAL 1	Calibrator 1	
CAL 2	Calibrator 2	
DISTRIBUTED IN THE USA BY	Distributed in the USA by	
DO NOT FREEZE	Do not freeze	
FOR USE WITH	Identifies products to be used together	
INFORMATION FOR USA ONLY	Information needed for United States of America only	
IVD	In Vitro Diagnostic Medical Device	
LOT	Batch code/Lot number	
MANUFACTURED FOR	Manufactured for	
PRODUCT OF JAPAN	Product of Japan	
REF	Catalog number/List number	
Rx ONLY	For use by or on the order of a physician only (applicable to USA classification only)	
$\triangle$	Caution	
l	Consult instructions for use	
	Manufacturer	
X	Temperature limitation	
$\Sigma$	Use by/Expiration date	

### Customer Service: Contact your local representative or find country-specific contact information on www.abbottdiagnostics.com.



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# HEMOGLOBIN A1c CONTROLS

This package insert contains information for the use of Hemoglobin A1° Controls on the ARCHITECT  $c\,8000$  and  $c\,4000$  Systems.

### Read Highlighted Changes: Revised April 2017.

Package insert instructions must be carefully followed. Reliability of assay results cannot be guaranteed if there are any deviations from the instructions in this package insert.

### INTENDED USE

For the estimation of test precision and the detection of systematic analytical deviations of the Hemoglobin  $A_{1c}$  assay on the ARCHITECT c 8000 and c 4000 Systems.

Refer to the Hemoglobin A1c reagent package insert and the ARCHITECT System Operations Manual for additional information.

### **CONTENTS / MATERIALS PROVIDED**

REF 4P52-11 Hemoglobin A1c Controls

- CONTROL L 1 x 1 mL
- CONTROL H 1 x 1 mL

Hemoglobin Ate Controls (lyophilized) contain hemoglobin and glycated hemoglobin from human whole blood. Prior to lyophilization, the control matrix is an MES-buffered solution. **CONTROL** and **CONTR** 

The value-assigned  $A_{1c}$  Control values are within the following hemoglobin  $A_{1c}$  ranges:

	CONTROL L	CONTROL H
Hemoglobin A1c Range	4.59 to 6.02 %HbA1c	9.42 to 11.07 %HbA1c

Actual analyte concentrations for this lot of controls are listed in the Hemoglobin  $A_{1c}$  Control value sheet, packaged with the controls. Verify that the lot number listed on each control vial agrees with the lot number printed on the value sheet.

### STANDARDIZATION

Each lot of controls is value-assigned. The glycated hemoglobin value in National Glycohemoglobin Standardization Program (NGSP) units (%HbA1c) and in International Federation of Clinical Chemistry and Laboratory Medicine (IFCC) units (mmol/mol HbA1c) are provided for each lot. Controls are prepared gravimetrically, lyophilized, and then value assigned using secondary calibrators that are traceable to the IFCC reference method.

### MATERIALS REQUIRED BUT NOT PROVIDED

- Calibrated pipette capable of measuring 1 mL
- Purified or deionized water
- Vortex (optional)

### PRECAUTIONS

### **Precautions for Users**

- · IVD
- For In Vitro Diagnostic Use.
- · Rx ONLY
- Do not use components beyond the expiration date.
- · Do not mix components from different kit lot numbers.



CAUTION: This product contains human-sourced and/or potentially infectious components. Refer to the CONTENTS / MATERIALS PROVIDED section of this package insert. No known test method can offer complete assurance that products derived from human sources or inactivated microorganisms will not transmit infection. Therefore, all human-sourced materials should be considered potentially infectious. It is recommended that these controls and human specimens be handled in accordance with the OSHA Standard on Bloodborne Pathogens.<sup>1</sup> Biosafety Level 2<sup>2</sup> or other appropriate biosafety practices<sup>3,4</sup> should be used for materials that contain or are suspected of containing infectious agents. The human blood used in the controls is nonreactive for HBsAg, HIV-1 RNA or HIV-1 Ag, anti-HCV, and anti-HIV-1/ HIV-2



The following warnings and precautions apply to: [CONTROL L] and [CONTROL H]

Harmful to aquatic life.

Causes serious eye irritation.

Wash hands thoroughly after handling. Avoid release to the environment.

Wear protective gloves / protective clothing /

	WARNING:
$\cdot$	H319
	H402*
	Prevention
	P264
	P273*
	P280
	Response

P501\*

	eye protection.
Response	
P305+P351+ P338	IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
P337+P313	If eye irritation persists: Get medical advice / attention.
Disposal	

Dispose of contents / container in accordance with local regulations.

# \* Not applicable where regulation EU 1272/2008 (CLP) has been implemented.

- Safety Data Sheets are available at www.abbottdiagnostics.com or contact your local representative.
- For a detailed discussion of safety precautions during system operation, refer to the **ARCHITECT System Operations Manual**, *Section 8.*

### STORAGE AND STABILITY

Unopened, lyophilized Hemoglobin  $A_{1c}$  **CONTROL** and **CONTROL** are stable until the expiration date when stored at 2 to 8°C.

Store reconstituted controls at 2 to 8°C and use within 10 days. Do not freeze.

### **PREPARATION OF CONTROLS**

- The lyophilized controls should be reconstituted as follows:
- 1. Add 1.0 mL of purified or deionized water.
- Close bottle and allow the contents of the bottle to sit at room temperature for at least 10 minutes.
- 3. Mix thoroughly by low speed vortexing or by gently inverting 10 times before use. Avoid the formation of foam.

NOTE: After reconstitution, the controls do not require pretreatment.

### INSTRUCTIONS FOR USE

The Hemoglobin  $A_{1\circ}$  assay uses the Hemoglobin  $A_{1\circ}$  Controls for the Hemolysate application.

To order the controls:

- Enter the control values into the control file from the Hemoglobin A<sub>1</sub>c Control value sheet.
- Mix controls several times by gently inverting to ensure homogeneity. Avoid the formation of foam.
- 3. Open the bottles, place an appropriate amount of each control into a separate sample cup.
- 4. Cap bottles and store at 2 to 8°C after use.
- Remove air bubbles, if present in the sample cup, with a new applicator stick. To minimize volume depletion, do not use a transfer pipette to remove the bubbles.

**CAUTION:** Bubbles in controls may interfere with proper detection of control level in the sample cup causing insufficient control aspiration that could impact results.

- 6. Place each sample cup in the assigned position.
- 7. For information on ordering controls, refer to *Section 5* of the **ARCHITECT System Operations Manual**.

### INDICATIONS OF INSTABILITY OR DETERIORATION

Instability or deterioration should be suspected if there are visible signs of leakage, extreme turbidity, microbial growth, or if control values do not meet the Hemoglobin A₁₀ Control value sheet and/or **ARCHITECT** System Operations Manual criteria.

### LIMITATIONS OF THE PROCEDURE

- The Hemoglobin  $A_{\rm 1c}$  Controls must only be used for the Hemolysate application of the Hemoglobin  $A_{\rm 1c}$  assay.
- This assay must be performed by qualified laboratory personnel, under appropriate laboratory conditions, solely for the intended use of the assay.

### **BIBLIOGRAPHY**

- 1. US Department of Labor, Occupational Safety and Health Administration. 29 CFR Part 1910.1030. Bloodborne Pathogens.
- US Department of Health and Human Services. *Biosafety in* Microbiological and Biomedical Laboratories, 5th ed. Washington, DC: US Government Printing Office, December 2009.
- 3. World Health Organization. *Laboratory Biosafety Manual*, 3rd ed. Geneva: World Health Organization, 2004.
- Clinical and Laboratory Standards Institute (CLSI). Protection of Laboratory Workers From Occupationally Acquired Infections; Approved Guideline—Fourth Edition. CLSI Document M29-A4. Wayne, PA: CLSI; 2014.

### TRADEMARKS

ARCHITECT, c8000, and c4000 are trademarks of Abbott Laboratories in various jurisdictions.

All other trademarks are property of their respective owners.

Key to Symbols		
-	-	
CONTROL	Control	
CONTROL H	High Control	
CONTROL L	Low Control	
DISTRIBUTED IN THE USA BY	Distributed in the USA by	
DO NOT FREEZE	Do not freeze	
FOR USE WITH	Identifies products to be used together	
IFCC RANGE	International Federation of Clinical Chemistry and Laboratory Medicine Range	
INFORMATION FOR USA ONLY	Information needed for United States of America only	
IVD	In Vitro Diagnostic Medical Device	
LOT	Batch code/Lot number	
MANUFACTURED FOR	Manufactured for	
NGSP RANGE	National Glycohemoglobin Standardization Program Range	
PRODUCT OF JAPAN	Product of Japan	
REF	Catalog number/List number	
Rx ONLY	For use by or on the order of a physician only (applicable to USA classification only)	
$\bigwedge$	Caution	
ī	Consult instructions for use	
	Manufacturer	
X	Temperature limitation	
$\Sigma$	Use by/Expiration date	

Customer Service: Contact your local representative or find country-specific contact information on www.abbottdiagnostics.com.



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# HEMOGLOBIN A1c

Package insert instructions must be carefully followed. Reliability of assay results cannot be guaranteed if there are any deviations from the instructions in this package insert.

WARNING: The Hemoglobin A1c assay has significant interference with the fetal hemoglobin (HbF). Hemoglobin A1c results are invalid for patients with abnormal amounts of HbF, including those with known Hereditary Persistence of Fetal Hemoglobin. For more information regarding the specific concentrations of HbF that were found to interfere with Hemoglobin A1c assay, refer to the SPECIFIC PERFORMANCE CHARACTERISTICS, Specificity section of this package insert.

### Read Highlighted Changes: Revised May 2018.

### **INTENDED USE**

The Hemoglobin A<sub>1</sub>c assay is used in clinical laboratories for the quantitative *in vitro* measurement of percent hemoglobin A<sub>1</sub>c (NGSP) or HbA<sub>1</sub>c fraction mmol/mol (IFCC) in human whole blood and hemolysate on the ARCHITECT *c* 8000 and *c* 4000 Systems.

Hemoglobin A1c measurements are used as an aid in the diagnosis of diabetes mellitus, as an aid to identify patients who may be at risk for developing diabetes mellitus, and for the monitoring of long-term blood glucose control in individuals with diabetes mellitus.

### SUMMARY AND EXPLANATION OF TEST

HbA1c is the fraction of hemoglobin A that is first reversibly, then irreversibly glycated at one or both *N*-terminal valines of the  $\beta$ -chain.<sup>1</sup> The longer red blood cells are in circulation and the higher the ambient glucose levels, the higher the concentration of HbA1c. HbA1c reflects the average blood glucose level during the preceding 2 to 3 months. The HbA1c assay is useful as an aid in the:

- diagnosis of diabetes mellitus,
- · identification of patients at risk for developing diabetes, and
- monitoring of patients with diabetes mellitus.<sup>2-6</sup>

For monitoring diabetic patients, it is recommended that glycemic goals are individualized following current professional society recommendations.<sup>7</sup> As recommended by the American Diabetes Association (ADA), patients in the range of 5.7-6.4 %HbA1c (39-46 mmol/mol) would be in the category of increased risk for diabetes and results  $\geq 6.5\%$  (48 mmol/mol) may aid in the diagnosis of diabetes.<sup>7</sup> Several studies, including the Diabetes Control and Complications Trial (DCCT), have shown that long-term control of diabetes can prevent complications such as cardiovascular disease, retinopathy, nephropathy, and neuropathy. Measurement of HbA1c can be invaluable in the monitoring of glycemic control of diabetic patients.<sup>8-10</sup> This method is certified by the National Glycohemoglobin Standardization Program (NGSP), standardized to International Federation of Clinical Chemistry and Laboratory Medicine (IFCC), and traceable to DCCT.

### PRINCIPLES OF PROCEDURE

The Hemoglobin A<sub>1c</sub> assay consists of two separate concentration measurements: glycated hemoglobin (HbA<sub>1c</sub>) and total hemoglobin (THb). The two concentrations are used to determine the percent HbA<sub>1c</sub> (NGSP units) or the hemoglobin fraction in mmol/mol (IFCC units). The individual concentration values of HbA<sub>1c</sub> and THb generated by the Hemoglobin A<sub>1c</sub> assay are used only for calculating the percent hemoglobin A<sub>1c</sub> or HbA<sub>1c</sub> fraction, and must not be used individually for diagnostic purposes.

The anticoagulated whole blood specimen is lysed automatically on the system for the Whole Blood application or may be lysed manually using the Hemoglobin A<sub>1c</sub> Diluent ( $\boxed{\texttt{A1cDIL}}$ ) for the Hemolysate application.

### Glycated Hemoglobin (HbA1c)

The Hemoglobin A1c assay utilizes an enzymatic method that specifically measures *N*-terminal fructosyl dipeptides of the  $\beta$ -chain of HbA1c.

- In the pretreatment process, the erythrocytes are lysed and the hemoglobin is transformed to methemoglobin by reaction with sodium nitrite.
- With the addition of Reagent 1 ( $[\underline{R1}]$ ) to the sample, the glycosylated *N*-terminal dipeptide (fructosyl-VH) of the  $\beta$ -chain of hemoglobin is cleaved by the action of protease. The hemoglobin is transformed to stable methemoglobin azide by the action of sodium azide and the concentration of the hemoglobin is determined by measuring absorbance.
- Addition of Reagent 2 (R2) starts a reaction and fructosyl peptide oxidase (FPOX) is allowed to react with fructosyl-VH. The HbA1c concentration is measured by determining the resultant hydrogen peroxide.

### Total Hemoglobin (THb)

The hemoglobin is oxidized to stable methemoglobin azide by the action of sodium nitrite and sodium azide and the concentration of the hemoglobin is determined by measuring absorbance (sample +  $\boxed{\textbf{R1}}$ ).

### Hemoglobin A1c Calculations<sup>11</sup>

The final result is expressed as %HbA1c (NGSP) or mmol/mol HbA1c (IFCC) and is automatically calculated by the system from the HbA1c/THb ratio as follows:



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**mmol/mol HbA**<sub>1c</sub> **IFCC:**  $HbA_{1c}$  (mmol/mol) = (HbA\_{1c}/THb

HbA1c (mmol/mol) = (HbA1c/THb)  $\times$  1000

%HbA1c DCCT/NGSP:

HbA1c (%) = IFCC x 0.09148 + 2.152 Methodology: Enzymatic

### REAGENTS

### **Reagent Kit**

REF 4P52-20 Hemoglobin A1c is supplied as a liquid, ready-to-use,

three-reagent kit which contains.	
R1 Reagent 1	1 x 52 mL
R2 Reagent 2	1 x 20 mL
A1cDIL Diluent	2 x 35 mL

Estimated tests per kit: 300

Calculation is based on the minimum reagent fill volume per kit.

Reactive Ingredients		Concentration
R1	10-(carboxymethylaminocarbonyl)- 3,7-bis(dimethylamino) phenothiazine sodium salt	0.000817%
	Protease (Bacterial)	< 1 MU/dL
R2	Peroxidase (Horseradish)	5 to 15 kU/dL
	Fructosyl-peptide-oxidase ( <i>E. coli,</i> recombinant)	300 to 900 U/dL
A1cDIL	Sodium nitrite	> 0.05 to < 0.3%

Inactive Ingredients: R1 contains sodium azide as a stabilizer and preservative. R1 and A1cDIL contain ProClin 300 as a preservative. R2 contains of loxacin as a preservative.

### REAGENT HANDLING AND STORAGE

### Reagent Handling

- R1, R2, and A1cDIL are ready for use.
- Remove air bubbles, if present in the reagent cartridge, with a new applicator stick. Alternatively, allow the reagent to sit at the appropriate storage temperature to allow the bubbles to dissipate. To minimize volume depletion, do not use a transfer pipette to remove the bubbles.

**CAUTION:** Reagent bubbles may interfere with proper detection of reagent level in the cartridge, causing insufficient reagent aspiration that could impact results.

### **Reagent Storage**

- Unopened reagents are stable until the expiration date when stored at 2 to 8°C.
- Reagent stability is 50 days (1,200 hours) if the reagent is uncapped and on board.
- When either the R1 or R2 reagent cartridge becomes empty, replace both cartridges.
  - A1cDIL can be replaced independently of R1 and R2

### Indications of Deterioration

Instability or deterioration should be suspected if there are visible signs of leakage, extreme turbidity, microbial growth, or if control values fall outside the acceptance criteria.

### WARNINGS AND PRECAUTIONS

- **Precautions for Use**
- IVD
- For In Vitro Diagnostic Use.
- · Rx ONLY
- Do not use components beyond the expiration date.
- Do not mix R1 and R2 from different kit lot numbers.
- Note: A1cDIL may be used with R1 and R2 from different kit lot numbers.

- WARNING: Do not use ARCHITECT sample cups for whole blood samples. Refer to the Assay Procedure section of this package insert for further information.
- CAUTION: This product requires the handling of human specimens. It is recommended that all human-sourced materials be considered potentially infectious and handled in accordance with the OSHA Standard on Bloodborne Pathogens.<sup>12</sup> Biosafety Level 2<sup>13</sup> or other appropriate biosafety practices<sup>14,15</sup> should be used for materials that contain or are suspected of containing infectious agents.
- The following warnings and precautions apply to: R1

101101	ing manninge	and procaditorio apply to. III	
	DANGER:	Contains N,N-dimethylformamide, methylisothiazolones, diethylenetriamine- pentaacetic acid, morpholinoethanesulfonic acid and sodium azide.	F
$\mathbf{A}$	H360	May damage fertility or the unborn child.	
! >	H317	May cause an allergic skin reaction.	
<u>、</u>	H316*	Causes mild skin irritation.	
•	EUH032	Contact with acids liberates very toxic gas.	•
	Prevention		
	P201	Obtain special instructions before use.	•
	P261	Avoid breathing mist / vapors / spray.	•
	P280	Wear protective gloves / protective clothing / eye protection.	
	P272	Contaminated work clothing should not be allowed out of the workplace.	
	Response		
	P302+P352	IF ON SKIN: Wash with plenty of water.	
	P333+P313	If skin irritation or rash occurs: Get medical advice / attention.	
	P362+P364	Take off contaminated clothing and wash it before reuse.	v
	P308+P313	IF exposed or concerned: Get medical advice / attention.	
	Disposal		
	P501	Dispose of contents / container in	

accordance with local regulations.

\* Not applicable where regulation EU 1272/2008 (CLP) or OSHA Hazard Communication 29CFR 1910.1200 (HCS) 2012 have been implemented.

### The following warnings and precautions apply to: R2

WARNING:	Contains citric acid.
H316*	Causes mild skin irritation.
P333+P313*	If skin irritation or rash occurs: Get medical
	advice / attention.

\* Not applicable where regulation EU 1272/2008 (CLP) or OSHA Hazard Communication 29CFR 1910.1200 (HCS) 2012 have been implemented.

### The following warnings and precautions apply to: DILUENT

	WARNING:	Contains maleic acid and methylisothiazolones and sodium nitrite.
•⁄	H317	May cause an allergic skin reaction.
•	H402*	Harmful to aquatic life.
	Prevention	
	P261	Avoid breathing mist / vapors / spray.
	P272	Contaminated work clothing should not be allowed out of the workplace.
	P273*	Avoid release to the environment.
	P280	Wear protective gloves / protective clothing / eye protection.
	Response	
	P302+P352	IF ON SKIN: Wash with plenty of water.
	P333+P313	If skin irritation or rash occurs: Get medical advice / attention.
	P362+P364	Take off contaminated clothing and wash it before reuse.
	Disposal	
	P501	Dispose of contents / container in accordance with local regulations.
lot apr	licable where	regulation EU 1272/2008 (CLP) or OSHA

\* Not applicable where regulation EU 1272/2008 (CLP) or OSHA Hazard Communication 29CFR 1910.1200 (HCS) 2012 have been implemented.

- Safety Data Sheets are available at www.abbottdiagnostics.com or contact your local representative.
- For a detailed discussion of safety precautions during system operation, refer to *Section 8* of the **ARCHITECT System Operations Manual**.

### SPECIMEN COLLECTION AND HANDLING

### **Suitable Specimens**

- Use only whole blood specimens collected by standard venipuncture techniques into plastic tubes. Acceptable anticoagulants are:
- Dipotassium EDTA
- Lithium heparin
- Sodium heparin
- Sodium fluoride/disodium EDTA
- Tripotassium EDTA

### **Preparation for Analysis**

- Follow the tube manufacturer's collection instructions for specimen collection tubes.
- Do not overfill specimen collection tubes. Whole blood samples greater than 78 mm in height from the bottom of tube will result in an instrument error and results will not be generated. Refer to *Section 10* of the **ARCHITECT System Operations Manual**.
- For testing whole blood samples less than 600 µL, use 12 x 75 mm polypropylene conical bottom tubes.
- · Do not centrifuge samples.
- Visually inspect the specimens. If fibrin clots or particulate matter is observed, remove with a clean applicator stick.
- Mix all specimens thoroughly by low speed vortexing or gently inverting 10 times prior to loading onto the ARCHITECT c System.
- Frozen specimens must be completely thawed prior to mixing.
   Visually inspect the specimens. If layering or stratification is observed, continue mixing until specimens are visibly homogeneous.

### **Specimen Storage**

Analyze fresh specimens if possible. If testing will be delayed, store specimens per the instructions below.

### Whole Blood

- Specimens may be stored for:
  - up to 8 hours at room temperature or
  - up to 7 days at 2 to 8°C
  - NOTE: Refer to the Suitable Specimens section of this package
  - insert. If testing will be delayed more than 7 days, store at -70°C or
  - colder. CAUTION: Whole blood specimens that require freezing must be stored at -70°C or colder.
- · Avoid more than one freeze/thaw cycle.
- NOTE: During storage, specimens may settle. It is good laboratory practice to mix specimens thoroughly prior to loading onto the system.

### Hemolysate

- Hemolyzed specimens may be stored for
  - up to 4 hours at room temperature or
  - up to 24 hours at 2 to 8°C.

NOTE: Refer to the Suitable Specimens section of this package insert.

- · Do not freeze hemolyzed specimens.
- NOTE: During storage, hemolyzed specimens may settle. It is good laboratory practice to mix specimens thoroughly prior to loading onto the system.

### PROCEDURE

### **Materials Provided**

REF 4P52-20 Hemoglobin A1c Reagent Kit

### Materials Required but not Provided

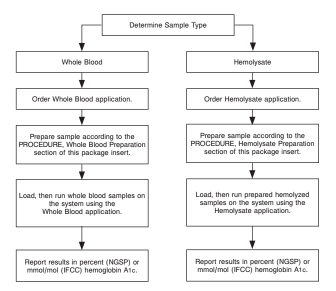
- REF 4P52-01 Hemoglobin A1c Calibrators
- REF 4P52-02 Hemoglobin A1c Calibrators The concentration of each calibrator is value-assigned and can change for each lot manufactured. Refer to the Hemoglobin A1c Calibrator Value Sheet.
- REF 4P52-10 Hemoglobin A1c Controls\*
- REF 4P52-11 Hemoglobin A1c Controls\* The concentration of each control is value-assigned with NGSP and IFCC values and can change for each lot manufactured. Refer to the Hemoglobin A1c Control Value Sheet.
- Commercially available whole blood controls\*\*
- 12 x 75 mm polypropylene conical bottom tubes\*\*
- Calibrated adjustable pipette capable of measuring 222 µL\*
- Calibrated micropipette capable of measuring 10 µL\*
- REF 01G48-04 c 4000/c 8000 Sample Probe
- Vortex (optional)
- \* Hemolysate application
- \*\* Whole Blood application

### **Assay Procedure**

- 1. Load the R1, R2, and A1cDIL on the ARCHITECT c8000 or c4000 System.
  - Place the A1cDIL in the reagent supply center (R1 c8000) for the Whole Blood application.
  - Configure the [A1cDIL] in the Configure Reagent Settings screen by selecting New, then Configure F6. Enter the reagent name as A1cDIL, the reagent type as Sample Diluent, the [A1cDIL] lot number and serial number as shown on the [A1cDIL] bottle label, and the R1 cartridge size as Small (55 mL cartridge). Select Add kit.
  - Assign the location of the <u>A1cDIL</u>. Refer to Load non-bar coded reagents in Section 5 of the ARCHITECT System Operations Manual.
- Perform a calibration for the Whole Blood and/or the Hemolysate application(s) as needed. Refer to Section 6 of the ARCHITECT System Operations Manual.
- 3. Refer to the flowchart below.

**NOTE:** No more than one replicate can be sampled from a sample cup or tube. To minimize the effects of evaporation, verify adequate sample volume is present before running the test.

For information on ordering patient samples and controls, refer to Section 5 of the **ARCHITECT System Operations Manual**.



Whole Blood Preparation

- · WARNING: Do not use ARCHITECT sample cups.
- WARNING: Do not overfill specimen collection tubes. Whole blood samples greater than 78 mm in height from the bottom of tube will result in an instrument error and results will not be generated. Refer to Section 10 of the ARCHITECT System Operations Manual.
- · Select the appropriate sample vessel using the table below:

If sample is:	Then:
	Do not use the Whole Blood application.
< 200 µL	Follow the instructions in the PROCEDURE, Hemolysate Preparation section of this package insert.
200 μL - 600 μL	Use 12 x 75 mm polypropylene conical bottom tubes only.
> 600 µL	Use suitable tubes or polypropylene conical bottom tubes.
> 78 mm in height in a large tube	Pipette 600 μL of sample into a suitable tube or polypropylene conical bottom tube.

### **Hemolysate Preparation**

- The minimum sample volume requirement is 150 µL for ARCHITECT sample cups.
- Prepare the hemolysate samples as follows:
- 1. Using a calibrated pipette, dispense 222  $\mu L$  <code>A1cDIL</code> into a tube or sample cup.
- 2. Using a calibrated micropipette, withdraw 10  $\mu L$  of the well-mixed whole blood patient specimen.
- 3. Wipe excess blood from the exterior of the pipette to ensure accurate transfer of the sample.

- Insert the pipette into the tube or sample cup containing the <u>A1cDIL</u> allowing the tip of the pipette to just make contact with the surface of the <u>A1cDIL</u> and dispense the 10 μL sample (1:23.2 dilution).
- Withdraw and dispense twice to rinse the pipette, always keeping the tip of the pipette in contact with the fluid in the tube.
- 6. Mix hemolysate thoroughly by low speed vortexing or by gently inverting 10 times. Avoid foaming.
- Allow the hemolysate to stand for a minimum of 1 minute at room temperature prior to testing.
- 8. If the hemolysate is prepared in a tube, transfer to a sample cup and place the cup on the instrument.

**NOTE:** The number of tests per kit is based on the 222  $\mu$ L of **A1cDIL** and 10  $\mu$ L of specimen volumes stated in steps 1 and 2 above. However, alternate volumes may be used for the 1:23.2 dilution, such as 555  $\mu$ L of **A1cDIL** and 25  $\mu$ L of specimen.

### **Specimen Dilution Procedure**

Specimens must not be diluted since the result is a calculated ratio. Refer to the RESULTS section of this package insert.

### CALIBRATION

- Calibration is stable for approximately 50 days (1,200 hours) and is required with each change in reagent lot number. Verify calibration with all levels of controls. If control results fall outside acceptable ranges, recalibration may be necessary.
- Both the Whole Blood and Hemolysate applications use the Hemoglobin A<sub>1</sub>c Calibrators (<u>REF</u>] 4P52-01 and 4P52-02), which are supplied separately.
- Hemoglobin A1c Calibrators are traceable to NGSP and IFCC reference methods.
- To perform a calibration, follow the instructions provided in the calibrator package insert.

### QUALITY CONTROL

The following is the recommendation of Abbott Laboratories for quality control. As appropriate, refer to your laboratory standard operating procedure(s) and/or quality assurance plan for additional quality control requirements and potential corrective actions.

Two levels of controls are to be run every 24 hours. The Hemoglobin A1c assay uses:

- Commercially available whole blood controls for the Whole Blood application. Follow the manufacturer's instructions for preparation of commercially available whole blood controls.
- Hemoglobin A1c Controls (REF) 4P52-10 and 4P52-11) for the Hemolysate application. Refer to the Hemoglobin A1c Control Value Sheet for NGSP and IFCC ranges.
- If more frequent control monitoring is required, follow the established quality control procedures for your laboratory.
- If quality control results do not meet the acceptance criteria defined by your laboratory, patient values may be suspect and corrective action should be taken. Follow the established quality control procedures for your laboratory. Recalibration may be necessary.
- Review quality control results and acceptance criteria following a change of reagent or calibrator lot.

### RESULTS

The individual concentrations of the Whole Blood and Hemolysate applications are measured by the system. NOTE: HbA1c or THb concentrations must not be used individually

for clinical purposes. IMPORTANT: Assay parameters must be configured exactly as defined in the ASSAY PARAMETERS section of this package insert.

 Refer to the LIMITATIONS OF THE PROCEDURE section of this package insert.

### **Conventional Units (NGSP)**

The percent HbA1 $\circ$  (%HbA1 $\circ$ ) is automatically calculated by the system per the calculation provided in the Hemoglobin A1 $\circ$  Calculations section. **SI Units (IFCC)** 

The hemoglobin A<sub>1c</sub> fraction (mmol/mol HbA<sub>1c</sub>) is automatically calculated by the system per the calculation provided in the Hemoglobin A<sub>1c</sub> Calculations section.

### LIMITATIONS OF THE PROCEDURE

- For use with ARCHITECT System software v8.1 or higher.
- The Hemoglobin A1c assay must not be used on the ARCHITECT c16000 System.
- This assay must be performed by qualified laboratory personnel, under appropriate laboratory conditions, solely for the intended use of the assay.

- · Do not centrifuge samples.
- · Do not freeze specimens that have been hemolyzed with the A1cDIL.
- Whole blood specimens that require freezing must be stored at -70°C or colder.
- Do not overfill specimen collection tubes. Whole blood samples greater than 78 mm in height from the bottom of tube will result in an instrument error and results will not be generated. Refer to Section 10 of the ARCHITECT System Operations Manual.
- Whole blood samples cannot be run in sample cups.
- Use specimen collection tubes, or for sample volumes < 600 µL use the 12 x 75 mm polypropylene conical bottom tubes as recommended in the PROCEDURE, Materials Required but not Provided section of this package insert.
- WARNING: The Hemoglobin A1c assay should not be used to diagnose diabetes during pregnancy. Hemoglobin A1c reflects the average blood glucose levels over the preceding 3 months (*i.e.*, the average life span of a red blood cell) and therefore may be falsely low during pregnancy or any other condition associated with recent onset of hyperglycemia and/or decreased red blood cell survival.<sup>16-19</sup>
- The Hemoglobin Atc assay should not be used to diagnose or monitor diabetes in patients with the following conditions:<sup>16-19</sup>
  - hemoglobinopathies except as demonstrated to produce acceptable performance (*e.g.*, sickle cell trait - refer to the SPECIFIC PERFORMANCE CHARACTERISTICS section of this package insert)
  - abnormal red blood cell turnover (*e.g.*, anemias from hemolysis and iron deficiency)
  - malignancies, and severe chronic hepatic and renal disease
- In cases of rapidly evolving Type 1 diabetes, the increase of HbA1c values might be delayed compared to the acute increase in glucose concentrations. In these conditions, diabetes mellitus must be diagnosed based on plasma glucose concentrations and/or the typical clinical symptoms.
- This test should not replace glucose testing for patients with Type 1 diabetes, pediatric patients, or pregnant women.
- The Hemoglobin A<sub>1c</sub> assay is susceptible to interference effects from conjugated bilirubin at > 15.0 mg/dL (180 μmol/L) and unconjugated bilirubin at > 10.0 mg/dL (171 μmol/L).
- The observed bias for samples containing HbC, HbD, HbE, HbS and HbA2 may be impacted by the method used to determine the reference Hemoglobin A1c concentration.
- Glycated HbF is not detected by the assay as it does not contain the  $\beta$ -chain that characterizes HbA1c. However, HbF is measured in the total Hb assay and as a consequence, specimens containing high amounts of HbF (> 5%) may result in lower than expected mmol/mol HbA1c values (IFCC) and %HbA1c values (NGSP).
- N-acetyl-4-benzoquinone imine (NAPQI), a metabolite of Acetaminophen at very high levels may lead to falsely low results.
- Dipyrone at high levels may lead to falsely low results.
- 4-methylamino antipyrine, a metabolite of Dipyrone at very high levels may lead to falsely low results.
- N-acetyl-L-cysteine at therapeutic levels may lead to falsely low results.
- Refer to the SPECIMEN COLLECTION AND HANDLING section of this package insert for specimen limitations.

### EXPECTED VALUES

For monitoring diabetic patients, it is recommended that glycemic goals are individualized following current professional society recommendations.<sup>7</sup> The American Diabetes Association (ADA) recommendations<sup>7</sup> are summarized in the following table.

HbA1c Value	Glycemic Goal
< 8 %HbA1c (64 mmol/mol)	Less stringent
< 7 %HbA1c (53 mmol/mol)	General (non-pregnant adults)
< 6.5 %HbA1c (48 mmol/mol)	More stringent

HbA1c values above 6.5 %HbA1c (48 mmol/mol) are an indication of hyperglycemia during the preceding 2 to 3 months or longer. According to the recommendations of the ADA, HbA1c values above 6.5 %HbA1c (48 mmol/mol) are suitable for the diagnosis of diabetes mellitus. Patients with HbA1c values in the range of 5.7 - 6.4 %HbA1c (39 - 46 mmol/mol) may be at a risk of developing diabetes.<sup>3,20</sup>

### SPECIFIC PERFORMANCE CHARACTERISTICS

### Measuring Interval

The measuring interval of the Hemoglobin A1c assay is 4.0 to 14.0 %HbA1c (20.22 to 129.51 mmol/mol HbA1c).

### Limit of Blank and Limit of Detection

Using a typical hemoglobin concentration of 8.2 mmol/L (13.2 g/dL) for the THb, the Hemoglobin A1c LoB result is 2.51 %HbA1c (3.89 mmol/mol) and LoD result is 2.52 %HbA1c (4.05 mmol/mol).

Limit of Blank (LoB) and Limit of Detection (LoD) were determined for each of the Hemoglobin A<sub>1</sub>c constituent assays (HbA<sub>1</sub>c and THb) using National Committee for Clinical Laboratory Standards (NCCLS) protocol EP17-A.<sup>21</sup>

The HbA<sub>1c</sub> constituent assay had an LoB of 31.9005  $\mu$ mol/L and an LoD of 33.2230  $\mu$ mol/L. The THb constituent assay had an LoB of 129.5129  $\mu$ mol/L and an LoD of 295.5947  $\mu$ mol/L.

### Linearity

Linearity was verified using NCCLS protocol EP6-A.22

### IFCC

The Hemoglobin A<sub>1c</sub> assay is linear across the range of 20.22 to 129.51 mmol/mol HbA<sub>1c</sub> based on an allowable tolerance of within or equal to  $\pm$  7%.

<u>SP</u>

The Hemoglobin A<sub>1</sub>c assay is linear across the range of 4.0 to 14.0 %HbA<sub>1</sub>c based on an allowable tolerance of within or equal to  $\pm$  5%. Specificity

### Hemoglobin Derivatives

A specificity study was conducted using Clinical and Laboratory Standards Institute (CLSI) protocol EP7-A2.<sup>23</sup> Specificity was assessed by comparing test samples containing the potential interferents listed below to reference samples. No interference was observed for:

- Acetylated Hemoglobin with  $\geq$  50 mg/dL of ASA (aspirin)
- Carbamylated Hemoglobin with ≥ 10 mmol/L of Cyanate
- Labile Hemoglobin with ≥ 1000 mg/dL of Glucose

### IFCC

The Hemoglobin A<sub>1c</sub> assay had a difference within  $\pm$  7% for samples with concentrations  $\geq$  38.78 mmol/mol HbA<sub>1c</sub>.

### NGSP

The Hemoglobin A<sub>1c</sub> assay had a difference within  $\pm$  5% for samples with concentrations  $\geq$  5.7 %HbA<sub>1c</sub>.

### Hemoglobin Variants

A specificity study was conducted using CLSI protocol EP7-A2.<sup>23</sup> Specificity was assessed by comparing the Hemoglobin A1c values to reference values for samples containing abnormal hemoglobins. Heterozygous hemoglobin variants (HbAS, HbAC, HbAD, HbAE, HbA2) do not interfere with the Hemoglobin A1c assay.

For the ARCHITECT c 8000 System, the data in %HbA1c (NGSP) are	
summarized in the following table.	

11	Relative % Difference from Reference Concentration					
Hemoglobin Variant	~ 6.0 %HbA1c	~ 9.0 %HbA1c				
HbC	-1.6	-1.9				
HbD	-0.8	1.8				
HbE	0.0	4.3				
HbS	-1.4	4.7				
HbA2	-0.6	-0.5				
HbF	Difference exceeds -5% when the amount of HbF in the sample exceeds 5% a					

<sup>a</sup> A negative % difference with HbF is proportional in magnitude to the % HbF present in the sample. For example, when the amount of HbF in the sample was 21.5%, the % difference was -18.5% on the ARCHITECT *c* 8000 System. Refer to the LIMITATIONS OF THE PROCEDURE section of this package insert for further information.

NOTE: The presence of multiple variants in a sample may impact the % difference.

For the ARCHITECT c4000 System, the data in %HbA1c (NGSP) are summarized in the following table.

	Relative % Difference from Reference Concentration							
	~ 6.0 %I (5.5 to 6.5		~ 9.0 %HbA1c (7.5 to 10.5 %HbA1c) <sup>a</sup>					
Hemoglobin Variant	Relative %			Range <sup>b</sup>				
HbC	-3.1	-6.9, 3.3	Difference -0.5	-4.2, 2.7				
HbD	0.6	-3.4, 3.2	0.2	-1.3, 2.6				
HbE	1.0	-3.3, 7.8	2.5	-2.1, 6.3				
HbS	-0.8	-3.6, 3.3	-0.5	-3.8, 2.2				
HbA2	0.7	0.0, 1.7	2.9	1.4, 4.5				
HbF	Difference exceeds -5% when the amount of HbF in the sample exceeds 5% °							

 $^a$  The HbA2 results at  $\sim 9.0$  %HbA1c consisted of samples between 7.2 to 11.2 %HbA1c.

 $^b$  The range is defined as the minimum and maximum relative % difference at each concentration level (~ 6.0 and ~ 9.0 % HbA1c).

<sup>c</sup> A negative % difference with HbF is proportional in magnitude to the % HbF present in the sample. For example, when the amount of HbF in the sample was 20.4%, the % difference was -20.0% on the ARCHITECT *c* 4000 System. Refer to the LIMITATIONS OF THE PROCEDURE section of this package insert for further information.

NOTE: The presence of multiple variants in a sample may impact the % difference.

### Interference

Interference studies were conducted using CLSI protocol EP7-A2.23 Interference effects were assessed by comparing test samples containing the potential interferents listed below to reference samples.

IFCC

The Hemoglobin A<sub>1c</sub> assay had a difference within  $\pm$  7% for samples with concentrations  $\geq$  38.78 mmol/mol HbA<sub>1c</sub>.

NGSP

The Hemoglobin A<sub>1</sub>c assay had a difference within  $\pm$  5% for samples with concentrations  $\geq$  5.7 %HbA<sub>1</sub>c.

For the ARCHITECT c 8000 and c 4000 Systems, the data in %HbA1c (NGSP) are summarized in the following tables.

	Interferent	Concentration	%	Interfe	erence	e <sup>a</sup>
			6.0 - 7.0 %HbA1c			8.0 bA1c
	Conventional	SI	ARC	HITEC	Г <i>с</i> Sy	stem
Potential Interferent	Units	Units	Ab	Bb	Ab	В <sup>b</sup>
Ascorbic Acid	3.0 mg/dL	0.15 mmol/L	0.0	0.0	0.0	0.0
Bilirubin (Conjugated) <sup>c</sup>	15.0 mg/dL	180 µmol/L	-3.2	-3.1	-2.2	-3.3
Bilirubin (Unconjugated) <sup>d</sup>	10.0 mg/dL	171 µmol/L	-3.0	-2.3	-2.2	-2.7
Glucose	1000 mg/dL	55.5 mmol/L	0.0	0.0	0.0	0.0
Rheumatoid Factor	200 IU/mL	200 IU/mL	0.0	0.0	0.0	0.0
Triglycerides	3000 mg/dL	33.9 mmol/L	-1.6	0.0	0.0	-4.5
Total Protein	22 g/dL <sup>e</sup>	220 g/L	0.0	0.0	0.0	-1.1
Urea	667 mg/dL	111.06 mmol/L	0.0	0.0	0.0	0.0
Vitamin E	8.6 mg/dL	200 µmol/L	0.0	1.6	0.0	0.0

Test Result - Control Result а % Interference = x 100 Control Result

<sup>b</sup> A = ARCHITECT c 8000 System; B = ARCHITECT c 4000 System

Samples containing conjugated bilirubin at > 15.0 mg/dL (180 µmol/L) demonstrated interference. Refer to the LIMITATIONS OF THE PROCEDURE section of this package insert for further information.

- <sup>d</sup> Samples containing unconjugated bilirubin at > 10.0 mg/dL (171 µmol/L) demonstrated interference. Refer to the LIMITATIONS OF THE PROCEDURE section of this package insert for further information.
- The total protein concentration of 22 g/dL includes serum protein as well as hemoalobin.

	Interferent C	Concentration	9	% Interfe	erencea	
			6.0 -		≥ 8	
			%Hb		%HbA1c	
Potential	Conventional	SI			Γ <i>c</i> Syst	r
Interferent	Units	Units	Ab	Bb	Ab	Bb
Acarbose	50 mg/dL	0.77 mmol/L	0.0	0.0	0.0	0.0
Acetaminophen	200 µg/mL	1324 µmol/L	-0.24	-0.59	-0.80	-0.63
N-acetyl-4- benzoquinone imine <sup>c</sup>	1 mg/L	6.71 μmol/L	-3.14	-3.26	-4.10	-4.35
N-acetyl-L- cysteine <sup>d</sup>	160 mg/L	981.6 µmol/L	-3.69	-3.26	-4.09	-3.06
Acetylsalicylate	50.8 mg/dL	2.82 mmol/L	0.0	0.0	0.0	0.0
Atorvastatin	0.06 mg/dL	600 μg Eq/L	0.0	1.6	0.0	0.0
Captopril	0.5 mg/dL	23 µmol/L	-1.5	0.0	-1.1	0.0
Chlorpropamide	74.7 mg/dL	2.7 mmol/L	0.0	0.0	0.0	-1.1
Cyanate	50 mg/dL	6.16 mmol/L	0.0	0.0	1.1	1.1
Dipyrone <sup>e</sup>	50 mg/L	150.2 µmol/L	-	-3.34	-	-4.18
ыруюпе	100 mg/L	300.3 µmol/L	-4.62	-	-4.91	-
Furosemide	6.0 mg/dL	181 µmol/L	0.0	0.0	0.0	1.1
Gemfibrozil	7.5 mg/dL	300 µmol/L	0.0	0.0	0.0	0.0
Ibuprofen	50 mg/dL	2425 µmol/L	0.0	0.0	0.0	1.1
Insulin	450 micro units per mL	450 micro units per mL	0.0	0.8	0.0	0.0
Losartan	5 mg/dL	0.11 mmol/L	0.0	0.0	0.0	0.0
Metformin	5.1 mg/dL	310 µmol/L	0.0	0.0	0.0	0.0
Nicotinic Acid	61 mg/dL	4.95 mmol/L	0.0	-1.5	0.0	-0.5
Propranolol	0.2 mg/dL	7.71 μmol/L	0.0	0.0	0.0	-0.5
Repaglinide	0.006 mg/dL	132.57 nmol/L	0.0	0.8	0.0	0.0
4-acetamido antipyrine	40 mg/L	163.3 µmol/L	-0.08	-0.11	0.09	0.00
4-aminoantipyrine	40 mg/L	197.0 µmol/L	-2.68	-2.47	-2.72	-2.49
4-formylamino antipyrine	40 mg/L	173.2 µmol/L	-0.09	-0.15	0.01	-0.05

4-methylamino antipyrine <sup>f</sup>	24 mg/L	110.6 µmol/L	-3.48	-4.43	-4.32	-4.07
a contractoria	Test Res	ult - Control Res	ult	100		

Control Result

- x 100

- A = ARCHITECT c 8000 System; B = ARCHITECT c 4000 System Samples containing N-acetyl-4-benzoquinone imine at > 1.0 mg/L
- (6.7 µmol/L) demonstrated interference.
- Samples containing N-acetyl-L-cysteine at > 160.0 mg/L (981.6 µmol/L) demonstrated interference.
- Samples containing Dipyrone at > 50.0 mg/L (150.2 µmol/L) demonstrated interference.
- Samples containing 4-methylamino antipyrine at > 24.0 mg/L (110.6 µmol/L) demonstrated interference.
- Refer to the LIMITATIONS OF THE PROCEDURE section of this package insert for further information.

### Precision

% Interference = -

A study was performed based on guidance from the NCCLS protocol EP5-A2.24 Testing was conducted using 3 lots of Hemoglobin A1c Reagents and Calibrators, 1 lot of commercially available controls, and 3 instruments. Two levels of controls and 3 levels of human whole blood panels were assayed in a minimum of 2 replicates at 2 separate times per day for 20 different days. Each reagent lot used a stored calibration curve throughout the study.

**IFCC** 

The Hemoglobin A1c assay is designed to have an imprecision of an SD of ≤ 1.42 mmol/mol HbA1c for samples with concentrations < 38.78 mmol/mol HbA1c,  $a \le 3\%$  within-laboratory (total) %CV for samples targeted to 47.53 mmol/mol HbA1c (38.78 to 53.00 mmol/mol HbA<sub>1c</sub>, inclusive), and  $a \le 5.0\%$  within-laboratory (total) %CV for samples with concentrations > 53.00 mmol/mol HbA1c. NGSP

The Hemoglobin A1c assay is designed to have an imprecision of an SD of  $\leq$  0.13 %HbA1c for samples with concentrations < 5.7 %HbA1c, a  $\leq$  2% within-laboratory (total) %CV for samples targeted to 6.5 %HbA1c (5.7 to 7.0 %HbA1c, inclusive), and a  $\leq$  3.5% within-laboratory (total) %CV for samples with concentrations > 7.0 %HbA1c. For the ARCHITECT *c* 8000 System, the data in %HbA1c (NGSP) are summarized in the following table.

						Within- Laboratory Precision		Component of	
	Instru-		Mean	Withi	n-Run	(Tot	al) <sup>a</sup>	Between-Lot	
Sample	ment	n	%HbA1c	SD	%CV	SD	%CV	SD	%CV
	1	240	4.9	0.01	0.3	0.02	0.5	0.03	0.5
Control 1	2	240	4.9	0.01	0.3	0.02	0.4	0.02	0.4
	3	240	4.8	0.02	0.4	0.03	0.6	0.03	0.6
	1	240	9.6	0.03	0.3	0.04	0.4	0.09	1.0
Control 2	2	239 <sup>c</sup>	9.6	0.03	0.3	0.04	0.4	0.08	0.9
	3	240	9.5	0.03	0.3	0.04	0.4	0.11	1.1
	1	240	4.4	0.01	0.3	0.02	0.4	0.03	0.7
Panel Near 4.0 %HbA1c	2	240	4.4	0.01	0.3	0.02	0.4	0.03	0.7
4.0 /010/10	3	240	4.4	0.02	0.3	0.02	0.5	0.03	0.6
	1	240	6.4	0.01	0.2	0.02	0.4	0.05	0.7
Panel Range 6.0 - 7.0 %HbA1c	2	240	6.4	0.02	0.3	0.02	0.3	0.04	0.6
0.0-7.0 /010/10	3	240	6.4	0.02	0.2	0.03	0.5	0.04	0.6
	1	240	8.9	0.02	0.2	0.03	0.3	0.06	0.7
Panel Range 8.0 - 10.0 %HbA1c	2	240	8.9	0.02	0.3	0.03	0.3	0.05	0.6
0.0-10.0 /01DATC	3	240	8.9	0.02	0.2	0.04	0.4	0.06	0.6

Contains within-run, within-day, and between-day variance components. Contains within-run, within-day, between-day, and between-lot variance

components.

A single replicate outlier with a %HbA1c value of 6.1 was removed from the analysis. The within-laboratory (total) %CV including the single replicate outlier was 2.4%

For the ARCHITECT c4000 System, the data in %HbA1c (NGSP) are summarized in the following table.

							hin- ratory	Precisio Addit	
	Instru-		Mean	Within-Run		Precision Within-Run (Total) <sup>a</sup>		Compoi Betwee	
Sample	ment	n	%HbA1c	SD	%CV	SD	%CV	SD	%CV
	1	240	4.9	0.01	0.3	0.02	0.5	0.02	0.5
Control 1	2	240	4.9	0.01	0.3	0.03	0.7	0.03	0.7
	3	240	4.9	0.01	0.2	0.02	0.4	0.03	0.6
	1	240	9.6	0.02	0.2	0.03	0.3	0.03	0.3
Control 2	2	240	9.6	0.02	0.2	0.03	0.4	0.04	0.4
	3	240	9.6	0.02	0.2	0.03	0.3	0.04	0.4

1	240	4.4	0.01	0.2	0.02	0.5	0.04	0.9
2	240	4.5	0.01	0.2	0.03	0.6	0.04	0.9
3	240	4.4	0.01	0.2	0.02	0.4	0.04	1.0
1	240	6.5	0.01	0.2	0.02	0.3	0.05	0.7
2	240	6.5	0.01	0.1	0.03	0.4	0.04	0.7
3	240	6.5	0.01	0.2	0.02	0.3	0.05	0.8
1	240	9.0	0.01	0.1	0.03	0.3	0.06	0.6
2	240	9.0	0.01	0.1	0.02	0.3	0.05	0.6
3	240	9.0	0.01	0.2	0.03	0.3	0.06	0.6
	3 1 2 3 1 2	2     240       3     240       1     240       2     240       3     240       1     240       2     240       3     240       1     240       2     240	2         240         4.5           3         240         4.4           1         240         6.5           2         240         6.5           3         240         6.5           1         240         9.0           2         240         9.0	2         240         4.5         0.01           3         240         4.4         0.01           1         240         6.5         0.01           2         240         6.5         0.01           3         240         6.5         0.01           3         240         6.5         0.01           3         240         6.5         0.01           3         240         9.0         0.01           1         240         9.0         0.01           2         240         9.0         0.01	2         240         4.5         0.01         0.2           3         240         4.4         0.01         0.2           1         240         6.5         0.01         0.2           2         240         6.5         0.01         0.1           3         240         6.5         0.01         0.1           3         240         6.5         0.01         0.1           3         240         6.5         0.01         0.1           3         240         9.0         0.01         0.1	2         240         4.5         0.01         0.2         0.03           3         240         4.4         0.01         0.2         0.02           1         240         6.5         0.01         0.2         0.02           2         240         6.5         0.01         0.1         0.03           3         240         6.5         0.01         0.1         0.03           3         240         6.5         0.01         0.1         0.03           3         240         6.5         0.01         0.1         0.02           1         240         9.0         0.01         0.1         0.03           2         240         9.0         0.01         0.1         0.03	2         240         4.5         0.01         0.2         0.03         0.6           3         240         4.4         0.01         0.2         0.02         0.4           1         240         6.5         0.01         0.2         0.02         0.3           2         240         6.5         0.01         0.1         0.03         0.4           3         240         6.5         0.01         0.1         0.03         0.4           3         240         6.5         0.01         0.1         0.03         0.4           3         240         6.5         0.01         0.2         0.02         0.3           1         240         9.0         0.01         0.2         0.02         0.3           1         240         9.0         0.01         0.1         0.03         0.3           2         240         9.0         0.01         0.1         0.02         0.3	2         240         4.5         0.01         0.2         0.03         0.6         0.04           3         240         4.4         0.01         0.2         0.02         0.4         0.04           1         240         6.5         0.01         0.2         0.02         0.3         0.05           2         240         6.5         0.01         0.2         0.02         0.3         0.04           3         240         6.5         0.01         0.1         0.03         0.4         0.04           3         240         6.5         0.01         0.1         0.03         0.4         0.04           3         240         6.5         0.01         0.2         0.02         0.3         0.05           1         240         9.0         0.01         0.1         0.03         0.3         0.06           2         240         9.0         0.01         0.1         0.02         0.3         0.05           2         240         9.0         0.01         0.1         0.02         0.3         0.05

<sup>b</sup> Contains within-run, within-day, between-day, and between-lot variance components.

### Method Comparison

The Hemoglobin A<sub>1c</sub> assay is designed to have a slope of 1.00  $\pm$  0.10 and a correlation coefficient (r) of  $\geq$  0.95 for specimens across the measuring interval when compared to an NGSP secondary reference laboratory method.

A correlation study was performed using CLSI protocol EP9-A2-IR<sup>25</sup> with Deming regression. Human whole blood specimen results from the Hemoglobin A1c assay were compared with those from an NGSP secondary reference laboratory method.

For the ARCHITECT *c* 8000 and *c* 4000 Systems, the data in %HbA1c are summarized in the following table.

	ARCHITECT vs.
	NGSP Secondary Reference Laboratory Method
Ν	128
Y - Intercept	-0.2
Correlation Coefficient	0.995
Slope	1.01
ARCHITECT Range (%HbA1c)	4.0 to 13.2 (ARCHITECT c 8000) 4.0 to 13.3 (ARCHITECT c 4000)
NGSP Secondary Reference Laboratory Range (%HbA1c)	4.0 to 13.4

### Bias

**IFCC** 

The Hemoglobin A<sub>1</sub>c assay is designed to have a bias of  $\leq$  5% at 42.06, 47.53, and 53.00 mmol/mol HbA<sub>1</sub>c using Deming regression. NGSP

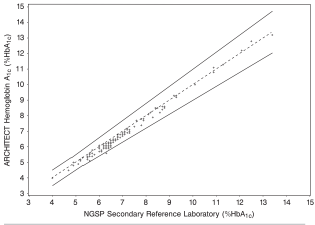
The Hemoglobin A<sub>1</sub>c assay is designed to have a bias of  $\leq$  3% at 6.0, 6.5, and 7.0 %HbA<sub>1</sub>c using Deming regression.

For the ARCHITECT *c* 8000 System, the bias in %HbA1c (NGSP) ranged from -3.0% to -2.4%. For the ARCHITECT *c* 4000 System, the bias in %HbA1c (NGSP) ranged from -2.3% to -1.8%.

### Allowable Total Difference (ATD) Zone

The Hemoglobin A<sub>1</sub>c assay is designed to have > 95% of observations in the ATD zone and the low limit of the two-sided 95% Confidence Interval (Cl) > 89.5% using Deming regression. For the ARCHITECT *c* 8000 System, the percentage of observations in

For the ARCHITECT *c* 8000 System, the percentage of observations in the ATD zone was 99.2% (127/128) and the lower limit of the two-sided 95% CI was 95.7%. The ATD zone plot is presented below.



For the ARCHITECT c 4000 System, the percentage of observations in the ATD zone was 100.0% (128/128) and the lower limit of the two-sided 95% CI was 97.1%.

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### TRADEMARKS

The ARCHITECT c System family of instruments consists of  $c\,4000,\,c\,8000,\,{\rm and}\,c\,16000$  instruments.

ARCHITECT, c 4000, c 8000, c 16000, and c System are trademarks of Abbott Laboratories in various jurisdictions.

All other trademarks are property of their respective owners.

### HbA1c Whole Blood—Conventional and SI Units

Configure assay paramete	rs — General							
		. <u>.</u>						
	O SmartWash O Res							
Assay: HbA1cWB	Type: Photometric	Version: †						
Number: 1106 Ass								
Run controls for onbo								
Reaction definition		O Validity checks						
Reaction mode: End		Read times						
	nary Secondary							
Wavelength: 660	/ 804	Main: 23 - 24						
Last required read: 24	00 0 0000 0-1	arraation.						
0	000 – 3.0000 Color o	Blank : <b>15 – 16</b>						
Sample blank type Sel		Biank : 15 - 16						
O Reaction definition	Reagent / Sample							
Beegente A1C00	Descent	=						
Reagent: A1C00 Diluent: A1cDIL	0	volume: 145 48 volume:						
Diluent dispense mode: Type 0		e mode: Type 0 Type 0						
Dilution nome Sample sam		Dilution Default factor dilution						
Dilution name Sample samp Std WB: 3.6 15.		1:23.22						
Std_WB: 3.6 15.	0 00	1.23.22						
O Reaction definition	O Reagent / Sample	Validity checks						
Beaction check: No		· validity checks						
Movin	Maximum alterations							
IVIAXIII	Maximum absorbance variation:							
Configure assay paramete								

Configure asso	ay parameters	- Calibiati			
O General	Calibration	O SmartWa	sh	O Results	O Interpretation
Assay: HbA1	SWB As	ssay number:	1106		
Calibration me	thod: Linear				
Calibrators	O Volu	mes	O Int	ervals	O Validity checks
Calibrator set:		Calibrator le	vel:		Concentration: ¥
HbA1c		Blank:	HbA1	c1	tt
		Cal 1:	HbA1	c2	tt
Replicates: 3	[Range 1 – 3]	Cal 2:	None		
		Cal 3:	None		
		Cal 4:	None		
		Cal 5:	None		
		Cal 6:	None		

O Calibrators	<ul> <li>Volumes</li> </ul>	01	ntervals	O Validit	y checks
Calibrator: HbA1c			Diluted		
	Calibrator level	Sample	sample	Diluent	Water
Blank:	HbA1c1	15.0			
Cal 1:	HbA1c2	15.0			
Cal 2:	None				
Cal 3:	None				
Cal 4:	None				
Cal 5:	None				
Cal 6:	None				

O Calibrators	O Volumes	Intervals	O Validity checks
Calibratio	n intervals:		
	Full interval: 1,200	(hours)	
Calibratio	n type:		
	Adjust type: None		

O Calibrators	tors O Volumes		Intervals	Validity checks
Blank	absorbance range:			
	Span:	Blank	<ul> <li>Blank</li> </ul>	
Span	Span absorbance range:		-	
E	xpected cal factor:	160.00		
Expected cal	factor tolerance %:	99		

Refer to System Configuration, Configuration Screen – Assay Settings in Section 2 of the ARCHITECT System Operations Manual for additional information.

- † Due to differences in instrument systems and unit configurations, version numbers may vary.
   Displays the number of decimal places defined in the decimal places parameter field.
- † Obtain value from Hemoglobin A1c Calibrator Value Sheet. Important: Value-assigned calibrator values are configured in the Calibrator set screen. It is also necessary to configure the Blank Concentration field in the assay parameter Calibration screen.
- \* Low-Linearity is the linear low value divided by the Standard dilution factor, then rounded up to the number of decimal places defined in the decimal places field.
- \*\* High-Linearity is set at the system maximum value. Since the Hemoglobin A1c is a ratio assay, the range of the assay is defined by the measuring interval.

Configure ass					
O General	O Calibration	SmartWash	O Resu	l <b>ts</b> O	Interpretation
Assay: HbA	1cWB				
COMPONENT	REAGENT / ASSAY	WASH		Volume	Replicates
R1	AMIK9	Detergent A	A	345	1
R1	DIG00	Detergent A	4	345	1
R1	DGT0B	Detergent A	4	345	1
R1	GENT9	Detergent A	4	345	1
R1	TOBRA	Detergent A	4	345	1
R1	VANCO	Detergent A	4	345	1
R1	TP000	0.5% Acid \	Wash	345	1
R2	AMIK9	Detergent A	4	345	1
R2	DIG00	Detergent A	4	345	1
R2	DGT0B	Detergent A	4	345	1
R2	GENT9	Detergent A	4	345	1
R2	TOBRA	Detergent A	4	345	1
R2	VANCO	Detergent A	4	345	1
Sample Probe*		Water			
* Sample Probe S	Sample wash protocol is	Maximum wa	ash.		

### HbA1c Whole Blood-Conventional and SI Units

Configure as	say parameter	s — Results		
O General	O Calibration	O SmartWash		
	Assay:	HbA1cWB		
	Assay number:	1106		
	Dilution default	range:	Result units:	umol/L
		Low-Linearity:	1.4308*	
		High-Linearity:	999999.9999**	
Gender and ag	e specific ranges:	- /		
GENDER	AGE (UNITS)	NORMAL	EXTREME	

Configure result units	
Assay:	HbA1cWB
Version:	t
Result units:	umol/L
Decimal places:	4 [Range 0-4]
Correlation factor:	1.0000
Intercept:	0.0000

### THb Whole Blood—Conventional and SI Units

• General         O Calibration         O SmartWash         O Results         O Interpretati           Assay:         THbWB         Type:         Photometric         Version:         †           Number:         1105         Assay availability:         Enabled         Photometric         Version:         †           Run controls for onboard reagents by:         Lot         Version:         •         •         •
Number:         1105         Assay availability:         Enabled           Run controls for onboard reagents by:         Lot         O         Validity checks
Run controls for onboard reagents by:         Lot           • Reaction definition         O Reagent / Sample         O Validity checks
Reaction definition     O Reagent / Sample     O Validity checks
Reaction mode: End up
Primary Secondary Read times
Wavelength: 476 / 804 Main: 15 - 16
Last required read: 16
Absorbance range: 0.0000 – 3.0000 Color correction: –
Sample blank type: None

O Reaction de	efinition		Reagent	/ Sample <sup>‡</sup>	01	alidity ch	ecks
						R1	R2
Reagent	t: A1C00			Reag	ent volume:	0	0
Diluent	t: <none></none>			Wa	ater volume:		
Diluent dispense	se mode: T	ype 0		Dispe	ense mode:	Type 0	Туре 0
Dilution name	Sample	Diluted sample		nt Water	Dilutic facto		Default dilution
::					=		٠

‡ Error code 0247 occurs when navigating from this screen to another. Select OK. Confirm that the empty assay parameter fields align with this insert. DO NOT enter any value. Select Cancel to exit the screen.

O Reaction defin		) Reagent /	Sample	Validity checks
Reaction ch	eck: None			
	Maximun	n absorbanc	e variation:	
Configure assa	ay parameters	— Calibra	tion	
O General	Calibration	O SmartWa	ish O Res	ults O Interpretation
Assay: THbV	VB Ass	ay number:	1105	
Calibration met	hod: Linear			
Calibrators	O Volum	es	O Intervals	O Validity checks
Calibrator set:		Calibrator le	vel:	Concentration: •
HbA1c		Blank:	HbA1c1	††
		Cal 1:	HbA1c2	<del>tt</del>
Replicates: 3	[Range 1 – 3]	Cal 2:	None	
		Cal 3:	None	
		Cal 4:	None	
		Cal 5:	None	
		Cal 6:	None	
○ Calibrators	Volum	85	∩ Intervals	O Validity checks

l	O Calibrato	rs	Volumes	01	ntervals	O Validit	y checks
	Calibrator:	HbA1c			Diluted		
l			Calibrator level	Sample	sample	Diluent	Water
		Blank:	HbA1c1	15.0			
		Cal 1:	HbA1c2	15.0			
		Cal 2:	None				
		Cal 3:	None				
		Cal 4:	None				
		Cal 5:	None				
		Cal 6:	None				

O Calibrators	O Volumes	Intervals	O Validity checks
Calibratio	n intervals:		
	Full interval: 1,200	(hours)	
Calibratio	n type:		
	Adjust type: None		

O Calibrators	O Volumes	0	In	tervals	۲	Validity checks
Blank	absorbance range:		-			
	Span:	Blank	-	Blank		
Span	absorbance range:		-			
E	xpected cal factor:	433.00				
Expected cal	factor tolerance %:	99				

Refer to System Configuration, Configuration Screen - Assay Settings in Section 2 of the ARCHITECT System Operations Manual for additional information.

† Due to differences in instrument systems and unit configurations, version numbers may vary.
 ♥ Displays the number of decimal places defined in the decimal places parameter field.

Displays the humber of decimal places defined in the decimal places parameter field.
† Obtain value from Hemoglobin A1c Calibrator Value Sheet. Important: Value-assigned calibrator values are configured in the Calibrator set screen.

It is also necessary to configure the Blank Concentration field in the assay parameter Calibration screen. \* Low-Linearity is the linear low value divided by the Standard dilution factor, then rounded up to the number of decimal places defined in the decimal

places field.

\*\* High-Linearity is set at the system maximum value. Since the Hemoglobin A1c is a ratio assay, the range of the assay is defined by the measuring interval.

O General	O Calibration	SmartWash	O Results	O Interpretation
Assay: TH	bWB			
COMPONENT	REAGENT / ASSAY	WASH	Volur	ne Replicates

### THb Whole Blood—Conventional and SI Units

Configure assay parameters — Results						
O General	O Calibration	O SmartWash				
	Assay:	THbWB				
	Assay number:	1105				
	Dilution default	range:	Result units:	umol/L		
		Low-Linearity:	12.7302*			
		High-Linearity:	999999.9999**			
Gender and age	e specific ranges:					
GENDER	AGE (UNITS)	NORMAL	EXTREME			

Configure result units	
Assay:	THbWB
Version:	†
Result units:	umol/L
Decimal places:	4 [Range 0-4]
Correlation factor:	1.0000
Intercept:	0.0000

Hb A1c Hemolysate—Conventional and SI Units
Configure assay parameters — General
General O Calibration O SmartWash O Results O Interpretation     Assay: HbA1cH Type: Photometric Version: †     Number: 1108 Assay availability: Enabled     Run controls for onboard reagents by: Lot
Reaction definition     O Reagent / Sample     O Validity checks
Reaction mode: End up Primary Secondary Read times Wavelength: 660 / 804 Main: 23 – 24 Last required read: 24
Absorbance range:     0.0000 - 3.0000     Color correction:        Sample blank type:     Self     Blank:     15 - 16
O Reaction definition • Reagent / Sample O Validity checks
Reagent:     A1C00     Reagent volume:     145     48       Diluent: <none>     Water volume:    </none>
Dilution name     Sample     Diluent     Water     factor     dilution       Std_H:     15.0       =     1:1.00     ●
O Reaction definition O Reagent / Sample ● Validity checks Reaction check: None
Maximum absorbance variation:
O General         ● Calibration         O SmartWash         O Results         O Interpretation           Assay:         Hb A1cH         Assay number:         1108           Calibration method:         Linear           ● Calibrators         O Volumes         O Intervals         O Validity checks
Calibrators O volumes O Intervals O validity checks
HbA1c         Blank:         HbA1c1
O Calibrators   Volumes O Intervals O Validity checks
Calibrator:       HbA1c       Diluted Calibrator level       Diluted sample       Diluted sample       Diluted         Blank:       HbA1c1       15.0
O Calibrators O Volumes ● Intervals O Validity checks Calibration intervals: Full interval: 1,200 (hours) Calibration type: Adjust type: None
O Calibrators O Volumes O Intervals ● Validity checks Blank absorbance range: Span: Blank - Blank Span absorbance range: Fynected cal factor: 160.00

Configure assay parameters — SmartWash					
O General	O Calibration	SmartWash O Re	sults (	O Interpretation	
Assay: Hb	A1cH				
COMPONENT	REAGENT / ASSAY	WASH	Volume	Replicates	
R1	AMIK9	Detergent A	345	1	
R1	DIG00	Detergent A	345	1	
R1	DGT0B	Detergent A	345	1	
R1	GENT9	Detergent A	345	1	
R1	TOBRA	Detergent A	345	1	
R1	VANCO	Detergent A	345	1	
R1	TP000	0.5% Acid Wash	345	1	
R2	AMIK9	Detergent A	345	1	
R2	DIG00	Detergent A	345	1	
R2	DGT0B	Detergent A	345	1	
R2	GENT9	Detergent A	345	1	
R2	TOBRA	Detergent A	345	1	
R2	VANCO	Detergent A	345	1	

### HbA1c Hemolysate—Conventional and SI Units

Configure assay parameters — Results					
O General	O Calibration	O SmartWash			
	Assay:	HbA1cH			
	Assay number:	1108			
	Dilution default	range:	Re	esult units:	umol/L
		Low-Linearity:	33.2230		
		High-Linearity:	999999.999	9**	
Gender and age specific ranges:					
GENDER	AGE (UNITS)	NORMAL		EXTREME	

Configure result units	
Assay:	HbA1cH
Version:	†
Result units:	umol/L
Decimal places:	4 [Range 0-4]
Correlation factor:	23.2000
Intercept:	0.0000

### Refer to System Configuration, Configuration Screen - Assay Settings in Section 2 of the ARCHITECT System Operations Manual for additional information.

† Due to differences in instrument systems and unit configurations, version numbers may vary.

• Displays the number of decimal places defined in the decimal places parameter field.

Expected cal factor: 160.00

99

Expected cal factor tolerance %:

- the Obtain value from Hemoglobin A1c Calibrator Value Sheet. Important: Value-assigned calibrator values are configured in the Calibrator set screen. It is also necessary to configure the Blank Concentration field in the assay parameter Calibration screen.
- \*\* High-Linearity is set at the system maximum value. Since the Hemoglobin A1c is a ratio assay, the range of the assay is defined by the measuring interval.

### THb Hemolysate—Conventional and SI Units

Configure assay parameters — General					
General O Calibrat	ion O SmartWash O Results	O Interpretation			
Assay: THbH	Type: Photometric Ve	ersion: †			
Number: 1107	Assay availability: Enabled				
Run controls for	onboard reagents by: Lot				
Reaction definition	O Reagent / Sample O Va	lidity checks			
Reaction mode:	End up				
	Primary Secondary	Read times			
Wavelength:	476 / 804 Main:	15 – 16			
Last required read:	16				
Absorbance range:	0.0000 – 3.0000 Color correction:				
Sample blank type:	None				
O Reaction definition	Reagent / Sample <sup>‡</sup> O Va	lidity checks			

						R1	R2
Reagent	: A1C00			Reagen	t volume:	0	0
Diluent	: <none></none>			Wate	r volume:		
Diluent dispense	se mode: T	ype 0		Dispens	se mode:	Type 0	Туре 0
Dilution name	Sample	Diluted sample	Diluent	Water	Dilutio factor		Default dilution
:				=			•

‡ Error code 0247 occurs when navigating from this screen to another. Select OK. Confirm that the empty assay parameter fields align with this insert. DO NOT enter any value. Select Cancel to exit the screen.

O Reaction definitio	n O Re	agent /	Samp	ole	Validity	v checks
Reaction check						
	Maximum ab	sorbanc	e varia	ation:		
Configure assay	parameters — (	Calibra	tion			
O General	Calibration O	SmartWa	ash	O Resul	ts Olm	terpretation
Assay: THbH		number:	1107	,		
Calibration method						
Calibrators			-	tervals		
Calibrator set:	Cal	ibrator le				centration: •
HbA1c		Blank:				_TT
Dealisation 0 1	Denne 1 01	Cal 1:			††	
Replicates: 3 [	Hange I – Sj	Cal 2: Cal 3:				
		Cal 3:		-		
		Cal 4.		-		
		Cal 6:		-		
				-		
O Calibrators	Volumes		O In	tervals	O Validit	y checks
Calibrator: HbA1c				Diluted		
	Calibrator level	· · · · ·	ole	sample	Diluent	Water
Blank:	HbA1c1	15.0				
Cal 1:		15.0				
Cal 2:						
Cal 3:						
Cal 4:						
Cal 5:						
Cal 6:	None					
O Calibrators	O Volumes	-	• Ir	ntervals	O Validit	v checks
Calibratio	n intervals:					
	Full interval: 1,20	0	(hou	rs)		
Calibratio						
	Adjust type: No	ne				

O Calibrators	O Volumes	0	In	tervals	Validit	y checks
Blank	absorbance range:		-			
	Span:	Blank	-	Blank		
Span	absorbance range:		-			
E	xpected cal factor:	433.00				
Expected cal	factor tolerance %:	99				

Refer to System Configuration, Configuration Screen - Assay Settings in Section 2 of the ARCHITECT System Operations Manual for additional information.

† Due to differences in instrument systems and unit configurations, version numbers may vary.

• Displays the number of decimal places defined in the decimal places parameter field.

the Obtain value from Hemoglobin A1c Calibrator Value Sheet. Important: Value-assigned calibrator values are configured in the Calibrator set screen. It is also necessary to configure the Blank Concentration field in the assay parameter Calibration screen. \*\* High-Linearity is set at the system maximum value. Since the Hemoglobin A1c is a ratio assay, the range of the assay is defined by the measuring interval.

O General	O Calibration	SmartWash	O Results	O Interpretation
Assay: TH				
COMPONENT	REAGENT / ASSAY	WASH	Volume	Replicates

### ThB Hemolysate—Conventional and SI Units

Configure assay parameters — Results							
O General	O Calibration	O SmartWash					
	Assay:	THbH					
	Assay number:	1107					
Dilution default range:			Result units:	umol/L			
		Low-Linearity:	295.5947	·			
		High-Linearity:	999999.9999**				
Gender and age specific ranges:							
GENDER	AGE (UNITS)	NORMAL	EXTREME				

Configure result units	
Assay:	THbH
Version:	†
Result units:	umol/L
Decimal places:	4 [Range 0-4]
Correlation factor:	23.2000
Intercept:	0.0000

### Percent A1c Whole Blood (NGSP)-Conventional Units

Configure assay parameters — General							
Gener	al O Calibration	O SmartWash	O Results	O Interpretation			
Assay:	%A1cWB	Type: Calo	culated				
Number:	3075 Ass	ay Availability: Ena	bled				
Fo	Formula: (ASSAY1/ASSAY2) * 1000 * 0.09148 + 2.152						
Select	ed assays:						
Minimum	Maximum						
ASSAY1:	HbA1cWB						
ASSAY2:	THbWB						
	·						

1c	Whole	Blood	(IFCC)—SI	Units

Α

Configure assay parameters — General						
General O Calibration O SmartWash O Results O Interpretation						
Assay:	A1cWB	Type: Calo	culated	-		
Number:	3074 As	say Availability: Ena	bled			
Fc	ormula: (ASSAY1/ASS	SAY2) * 1000				
Select	ed assays:					
Minimum	Maximum					
ASSAY1:	HbA1cWB					
ASSAY2:	THbWB					

Configure assay parameters — Results						
O General	O Calibration	O SmartWash	Results	O Interpretation		
	Assay:	%A1cWB				
	Assay number:	3075				
			Result u	nits: %		
Gender and ag	e specific ranges:					
GENDER	AGE (UNITS)	NORMAL	EXTREME			
	. ,					

Configure result units — Assay Settings				
Assay:	%A1cWB			
Version:	†			
Result units:	%			
Decimal places:	1 {Range 0 - 4]			

Configure assay parameters — Results						
O General	O Calibration	O SmartWash	Results	O Int	erpretation	
	Assay:	A1cWB				
	Assay number:	3074				
			Result u	inits:	mM/mol	
Gender and ag	e specific ranges:					
GENDER	AGE (UNITS)	NORMAL	EXTREME			
	. ,					

Configure result units — Assay Settings		
Assay:	A1cWB	
Version:	†	
Result units:	mM/mol	
Decimal places:	2 {Range 0 - 4]	

### Percent A1c Hemolysate (NGSP)-Conventional Units

Configur	Configure assay parameters — General			
Generation	al O Calibration	O SmartWash	O Results	O Interpretation
Assay:	%A1cH	Type: Ca	culated	
Number:	3077 Ass	ay Availability: Ena	abled	
Fo	rmula: (ASSAY1/ASS	AY2) * 1000 * 0.09	148 + 2.152	
Select	ed assays:			
Minimum	Maximum			
ASSAY1:	Hb A1cH			
ASSAY2:	THbH			

Configure assay parameters — Results				
O General	O Calibration	O SmartWash	Results	O Interpretation
	Assay:	%A1cH		
	Assay number: 3	3077		
			Result u	inits: %
Gender and age specific ranges:				
GENDER	AGE (UNITS)	NORMAL	EXTREME	
	. ,			

Configure result units — Assay Settings		
Assay:	%A1cH	
Version:	†	
Result units:	%	
Decimal places:	1 {Range 0 - 4]	

### A1c Hemolysate (IFCC)-SI Units

Configu	Configure assay parameters — General			
Gener	al O Calibratio	on O SmartWash	O Results	O Interpretation
Assay:	A1cH	Type: Cal	culated	
Number:	3076	Assay Availability: Ena	bled	
F	ormula: (ASSAY1/A	SSAY2) * 1000		
Select	ted assays:			
Minimum	Maximum			
ASSAY1:	Hb A1cH			
ASSAY2:	ТНЬН			

Configure assay parameters — Results					
O General	O Calibration	O SmartWash	Results	O In	terpretation
		1cH			
	Assay number: 3	076			
			Result	units:	mM/mol
Gender and age specific ranges:					
GENDER	AGE (UNITS)	NORMAL	EXTREME		

Configure result units — Assay Settings		
Assay:	A1cH	
Version:	†	
Result units:	mM/mol	
Decimal places:	2 {Range 0 - 4]	

Refer to **Assay Configuration** in *Section 2* of the **ARCHITECT System Operations Manual** for additional information. † Due to differences in instrument systems and unit configurations, version numbers may vary.

Key to Symbols		
A1cDIL	Hemoglobin A1c Diluent	
CONTAINS: AZIDE	Contains sodium azide. Contact with acids liberates very toxic gas.	
DISTRIBUTED IN THE USA BY	Distributed in the USA by	
FOR USE WITH	Identifies products to be used together	
INFORMATION FOR USA ONLY	Information needed for United States of America only	
IVD	In Vitro Diagnostic Medical Device	
LOT	Batch code/Lot number	
MANUFACTURED FOR	Manufactured for	
PRODUCT OF JAPAN	Product of Japan	
R1	Reagent 1	
R2	Reagent 2	
REF	Catalog number/List number	
Rx ONLY	For use by or on the order of a physician only (applicable to USA classification only)	
SN	Serial number	
i	Consult instructions for use	
	Manufacturer	
Σ	Sufficient for	
l X	Temperature limitation	
	Use by/Expiration date	

Customer Service: Contact your local representative or find country-specific contact information on www.abbottdiagnostics.com.



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# A Promise for Life

This document certifies that: Sergiu Sorocovici

has completed

# Architect i2000SR

Level1 / Level 2 Application, Operation, Troubleshooting from 9 February 2015 to 13 February 2015

Trainer : Athanasios Plakas

Date: 13 Feb 2015