Operation Manual Version 2.0

MR9610, SmartReader™ UV-Vis Microplate Reader





This instrument is for research use only



Foreword

Thank you for purchasing our MR9610, SmartReader™ UV-Vis Microplate Reader. This user manual describes how the instrument works and includes an operation guide, please read it carefully before operation and keep for future reference.

Initial check

Please check the instruments as well as all accessories with the packing list when first received and unpackaged. If you find any missing, please contact your distributor or Accuris Instruments.

Safety Warning and Guidelines

1. Important information for safe use

Users should have a clear understanding on use this instrument before operation. Please read this user manual carefully.



Please read the manual carefully and operate safely according to all guidelines.



This instrument is intended for Research Use Only

2. Safety Tips

The operation, maintenance and repair of the Instrument should comply with the basic guidelines and the warnings below.



Indoor use only.



Warning: Biological contamination hazard. All samples for testing, quality control, and calibration should be regarded as infectious, and any parts that come into contact with samples should also be treated as infectious. Please wear gloves and appropriate safety gear when operating this device.



Before using the device, read the operation manual carefully. These instruments are designed for use in a laboratory environment. The device must be operated by skilled laboratory personnel with the appropriate training.



Warning: Avoid injury. Keep your body or any part of body at a safe distance (1 foot or more) from the instrument when running.



The operator should not open or perform any repairs on the Instrument without specific instructions from Accuris Instruments or an authorized agent of Accuris. Performing any unauthorized service repairs will result in loss of warranty coverage. Contact Accuris Instruments or an authorized distributor or agent in the event of any service issues.



Before powering on, confirm that the voltage used meets the electrical requirements of the instrument as stated on the rating plate. If the electric cord is damaged, replace it with the same type. Hold the socket firmly when you pull the plug from an outlet, and don't pull the electric line only.



The Instrument should be installed in an environment of standard room temperature, low dust, low humidity and away from direct sunlight. Also make sure there is adequate ventilation, no corrosive gas or strong magnetic field.



Always power off the instrument when you are finished using it. Unplug the power cord and cover the instrument with a cloth or plastic sheet to prevent excessing dust from entering the housing.

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Pull the connector plug from the electrical outlet immediately and contact the vendor in the case of:

- Liquid entering the housing.
- Abnormal operation: such as any abnormal sound or smell.
- The instrument is dropped or there is any damage to the housing.
- Any malfunction

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Chapter 1 Introduction

The SmartReader[™] UV-Vis is an automated microplate reader for ELISA assays. Through absorbance readings, it can determine the concentration, as well as positive or negative detection of antibody and antigen in samples when used with ELISA kits and solutions. The instrument can also be used to test for purity and quantitate nucleic acids and proteins.

Key Features:

- 1) 10-inch color touch screen.
- 2) Operating system allows acquisition, editing and saving of data.
- 3) Stand-alone system incorporates built-in processing, it can be optionally connected to a PC with the use of SmartReader software.
- Graphical interface displays a representative plate for easy setting of well parameters: blank, sample, standards, positive/negative, quality control and multi-value comparisons.
- 5) Dual optical channels, one used for sample detection, and one used for reference.
- 6) End-point method, kinetic readings, and spectral scanning can be performed.
- 7) Accepts plates with or without lids. Top heating prevents condensation.
- Xenon lamp offers long life: 10⁹ flashes (1,000,000,000), equivalent to approximately 1,300,000 plate reads.
- Plate incubation feature allows accurate and uniform temperature control (≤0.5°C deviation between wells).
- 10) Top reading optical path with continual self-checking and automatic calibration option for monochromator wavelength and plate positioning.
- 11) Shaking function, time and speed are adjustable.
- 12) USB flash drive data export, for fast and easy data transfer.
- 13) Multi-user operation, data management for separate users.

Chapter 2 Features

Working conditions:

Ambient temperature: 10°C~40°C

Relative humidity: 30%~80% (Non-condensing environment)

Power: AC 100-240V 50-60Hz 2A

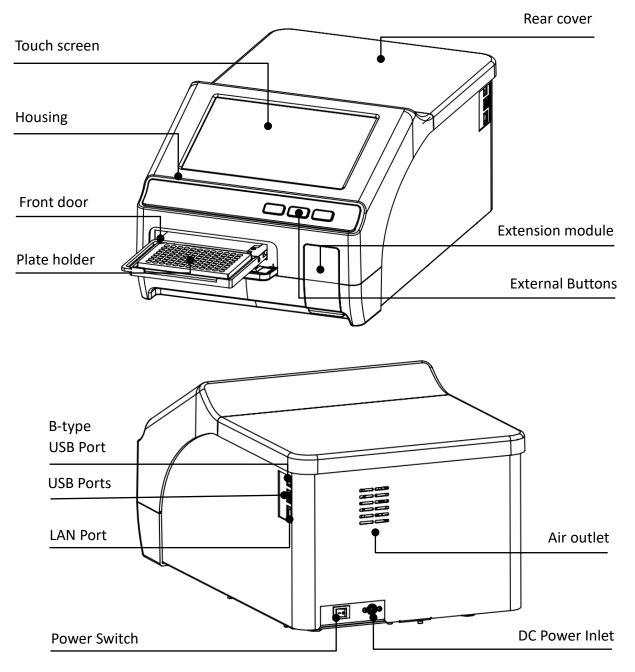
Parameters:

Model Parameter	MR9610
Light source	Xenon flash lamp >10 ⁹ flashes
Wavelength	200~1000nm
Bandwidth	< 2.5nm
Wavelength accuracy	±2nm
Wavelength repeatability	0.2nm
Read-out range	0.0-4.0 OD
Linearity@450nm	$R^2 \ge 0.999$, [0.0 – 3.0]
Accuracy@450nm	1.0% + 0.003 Abs (0 - 2.0] 2.0% (2.0 - 2.5 Abs]
Precision@450nm	CV < 0.5% or SD<0.003 (Precision mode); CV < 1.0% (Fast mode)
Stability@450nm	< 0.005A , (0.0 - 2.0] < 2% , (2.0 - 2.5]
Measurement speed	< 8 seconds in Fast mode (96-well plate) < 28 seconds in Precision mode (96-well plate) < 74 seconds in Precision mode (384-well plate)
Plate shaking	Linear (Slow : 16.8Hz, 1mm; Medium : 6.68Hz, 3mm; High : 2.44Hz, 15mm)
Incubation range	RT+ 5°C to 45°C
Temp. uniformity	± 0.5°C @ 37°C

Connections	 B-type USB port for PC connection Ethernet port A-type of USB ports for mouse and printer connections
Power requirements	Included power adapter: DC24V 6.67A 160W, AC 100V – 240V
Dimension (W \times D \times H)	$300{\times}500{\times}260mm$ / 11.8 x 19.7 x 10.25 inch
Weight (kg)	15.5kg / 34 Lbs

Chapter 3 Instrument Overview

Structure



Chapter 4 Installation

1. Initial check

Each MR9610 is thoroughly tested before shipping, but please check it again when you receive the instrument and contact your local distributor or Accuris Instruments if:

- The outer package is damaged
- The outer package has any obvious moisture stains
- The outer package has marks of impact

After opening, please check the instrument and box contents.

Confirm that all ordered accessories have been included.

Check the instrument's appearance for any damage.

2. Installation location and environment

• Installation location: locate the instrument on a flat, dry, clean and stable surface, allow enough room on the front side so there is space for plate holder drawer to move in and out, and also keep 15cm / 6 inches of space behind and on the sides for any wires and air flow.

- Environment:
 - a. The environment for installation should not be excessively humid or dusty.
 - b. Temperature should be within the range of $+10^{\circ}C \sim +40^{\circ}C$.

c. Relative humidity should be within the range of 30% ~ 80% to avoid condensation. Note: KEEP THE INSTRUMENT AWAY FROM CORROSIVE GASSES AND LIQUIDS.

3. Installation steps

① Place the instrument on a stable and level surface.

Note: DO NOT loosen any screws or parts without first consulting with the Accuris Instruments Service Department or your authorized distributor. Doing so can cause damage and void the warranty.

2 Connect the instrument to an appropriate power outlet using the provided power cord.

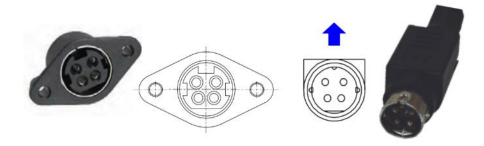


Fig. 1

Note: One cord coming from the power adapter will fit to the input jack on the back of the instrument. Make sure to connect it according to the above picture. The other cord will connect to a standard electrical outlet (100VAC to 230VAC)

③ After connecting the power cords, switch the power button on the rear of the instrument to the ON position, the front panel will cycle through a start-up and self-test screen.

Warning: Do not connect the instrument to a power outlet that is not properly grounded.

Chapter 5 Operation Guide

1. Initial self-check

This chapter explains a standard protocol set up and operation, beginning with the automatic self-check after power on. The following screen image will appear during the self-check:



Fig 2

The user login interface will appear after the self-check, see Fig 3. This allows login as the Admin, existing user, or click "Guest".

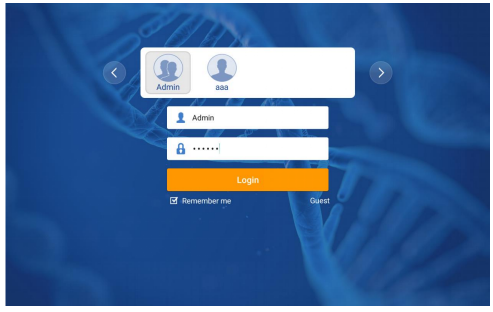


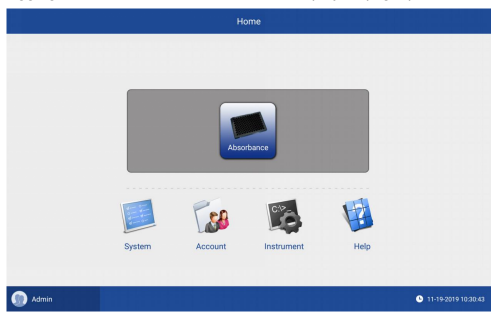
Fig 3

— 7 —

User type	Creation Method	Default password	Permission	Export
Admin	Cannot be deleted	"123456"	For all files of Admin, User and Guest	All can be
User	Created by Admin	Default is "123456" or is set when creating	Access to files under the user	exported
Guest	Cannot be deleted	No password	account	

Table 1

Note: Please keep the Admin password recorded and in a safe place, and contact Accuris Instruments or your distributor if the Admin password is lost or forgotten.



After logging in, the Home Interface screen is displayed (Fig 4.)

Fig 4

The "Admin" button on the lower left is for the logout and login interface.

2. System settings

In the "Systems" section, users create system settings (time, date and language) according to their needs, for details see Fig 5.

Chapter 5 Operation Guide

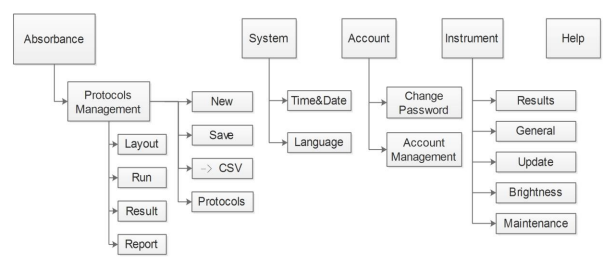


Fig 5

Note: 1. The instrument must be manually powered off and back on after the date and time settings are changed.

2. The Maintenance function is accessible only by authorized service representatives.

3. Click the "Home" button on the top left corner to go to the main interface.

3. Protocol Management

Click "Absorbance" on the home interface to enter the layout interface (fig 6). This interface is composed of six sections: the navigation bar at the top, the sidebar, the main display area, the optional bar, setting type selection and the status bar at the lower right area. The SmartReader UV-Vis incorporates a touch screen control panel, but the included mouse can also be used for all programming functions.

Home	Protocol 2019	Name 91119_	.10321	15			Navi	gatio	n ba	r				Image: Contract of the second sec
												Reset	List	Pipette Content
Layout		1	2	3	4	5	6	7	8	9	10	11	12	Clear the Type
	А													
Run	В													
Result	С													
	D													
Report	E				ſ	Main	disp	olay a	area					
керон	F													The optional bar
	G													
글	н													
The Sidebar	ſ	Select Blank	Stan	dard	Quality	Settin;	g Typ ^{known}	e Sel		1 Negative		Clear		
														11-19-2019 10:32:21

Fig 6

Protocol Name 20191119_103215

Protocol name can be modified by clicking it directly.



The default name of a new protocol is the system date and time, and it can be modified manually. A pop-up window will appear to remind the user when a modified protocol has not been saved.



Save the current protocol to the protocol list.



Shortcut for exporting raw data to a flash drive in .csv format. (Exports only the data displayed in the "Main Display Area")



Open the Protocol folder for accessing saved protocols, sorting, deleting, importing, exporting, and renaming protocols. The Protocol list opens on the right side of the screen and can be closed by clicking on the left area.

Acc	ount Ad	min	Edit		Adr	nin	Cano
	Q s	Search			Name	Date 🔻	Тур
	Name	Date 🔻	Туре		demo1	2022/01/18 04:19:11	Lock
	demo1	2022/01/18 04:19:11	Locked		demo2	2009/09/01 00:00:28	Lock
	demo2	2009/09/01 00:00:28	Locked		demo3	2022/01/18 04:17:11	Lock
	demo3	2022/01/18 04:17:11	Locked		2022127new	2022/01/28 05:06:24	Dat
	2022127new	2022/01/28 05:06:24	Data		20220119_044133	2022/01/19 23:17:20	Dat
	20220119_044133	2022/01/19 23:17:20	Data		20220119_041650	2022/01/19 04:18:05	Dat
	20220119_041650	2022/01/19 04:18:05	Data		20220119_003437	2022/01/19 00:34:53	Dat
	20220119_003437	2022/01/19 00:34:53	Data				
	Fotal : 7 Files				Selected : 0 Files		
Ł	Import 🕎 I	Rename 避	Save as	1	Delete	L	Ехро



When the protocol list is in the standard state, users can perform the below operations:

- Search: Enter a keyword to search the list.
- Sort: protocols can be sorted according to "Name", "Date" and "State". "demo1", "demo2" and "demo3" are always displayed at the first three positions.
- Import: For importing protocols from a flash drive to the instrument.
- Rename: Change the name of a protocol.
- Save as: Save as a new protocol.
- Edit: "Edit" button is located on the top right corner, see Fig 7 to enter the edit state.
- Account: Protocols of other accounts can be checked, but this function is only available for the Admin account.

When the protocol list in the edit state, the below operations are available:

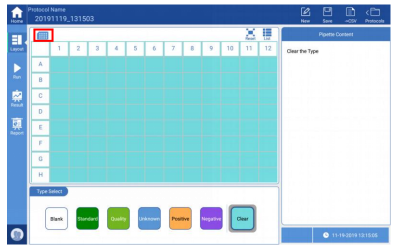
- Sorting: protocols can be sorted according to "Name", "Date" and "State".
- Checkbox: " \Box " can used to select more than one for batch operation.
- Delete: delete selected protocols.
- Export: For exporting selected protocols to a flash drive.
- Cancel: Return to the standard state of the protocol list.

4. Reading a Microplate

After a protocol name is created, the next step is to set the parameters according to the experiment requirements.

4.1 Plate layout settings

The interface will automatically display the plate in a 96 well plate layout when creating a new protocol. To change from a 96 to 384 well-layout, click on the top left of the plate layout to choose the required layout format.





The plate layout menu allows choosing 6 well-setting options, as well as a clear option. Choose the appropriate setting type, and the option bar will change accordingly, then click the corresponding wells on the main display area to finish the settings.

Note: the entire plate (all wells) can be selected by clicking the blank box at the upper left corner of the plate layout.



Blank: Set for a blank control during measurement. The average absorbance value of the blank wells is subtracted from the absorbance values of the other wells after a plate read.

Standard: Used for creating the standard curve. After clicking "Standard", the option bar will allow setting replicates and also calculations for quickly setting the standard concentrations, see Fig 9.

- Replicates: Use to set the same standard for multiple wells.
- Concentrations: After setting the concentration and unit of the first well, the protocol will calculate and set subsequent standards automatically according to the set operator and step size, or the values can be modified manually by clicking the corresponding wells.

Example: If the set concentration is $125 \text{ ng/}\mu\text{L}$, the operator is "×" (multiply) and the step size is "2", the concentration of the first well will be $125 \text{ ng/}\mu\text{L}$, the second well $250 \text{ ng/}\mu\text{L}$ (2 x $125 \text{ ng/}\mu\text{I}$) and the third $500 \text{ ng/}\mu\text{L}$ (2 x $250 \text{ ng/}\mu\text{L}$)... etc.

Quality

Quality control. Settings are done in the same manner as the wells of standard samples.

Unknown samples. Users can set wells as unknown, and there in an additional "Factor" option if the unknown sample has been diluted. If a dilution has been used, set the value for the dilution factor "X. 1:X means the unknown sample was diluted X times.



Positive Control, users can set several wells as positive controls



Clear

Negative Control, users can set several wells as negative controls

Used for clearing the settings of the plate layout.

Standard

f Home	Protocol 2019	Name 91119	_1315	03										New	C Save	.⇒csv	< 📩 Protocols
												Reset	List		Pipette Co	ontent	
Layout		1	2	3	4	5	6	7	8	9	10	11	12	Standard		34	
	Α	Std 01 1.000 Group 1	Std 01 1.000 Group 1	Std 01 1.000 Group 1	Un 01 1:1.00 Group 1	Un 05 1:1.00 Group 1	Un 09 1:1.00 Group 1	Std 10 30.000 Group 1	Std 11 60.000 Group 1	Std 12 120.000 Group 1	Un 13 1:1.00 Group 1	Un 17 1:1.00 Group 1	Un 21 1:1.00 Group 1	Replicates			\bigcirc
Run	В	Std 02 2.000 Group 1	Std 02 2.000 Group 1	Std 02 2.000 Group 1	Un 02 1:1.00 Group 1	Un 06 1:1.00 Group 1	Un 10 1:1.00 Group 1	Std 13 240.000 Group 1	Std 14 480.000 Group 1	Std 15 960.000 Group 1	Un 14 1:1.00 Group 1	Un 18 1:1.00 Group 1	Un 22 1:1.00 Group 1	Concentration	s		
~	С	Std 03 3.000 Group 1	Std 03 3.000 Group 1	Std 03 3.000 Group 1	Un 03 1:1.00 Group 1	Un 07 1:1.00 Group 1	Un 11 1:1.00 Group 1	Std 16 1920.000 Group 1	Std 17 3840.000 Group 1	Std 18 7680.000 Group 1	Un 15 1:1.00 Group 1	Un 19 1:1.00 Group 1	Un 23 1:1.00 Group 1	Conc.		1.	000
Result	D	Std 04 4.000 Group 1	Std 04 4.000 Group 1	Std 04 4.000 Group 1	Un 04 1:1.00 Group 1	Un 08 1:1.00 Group 1	Un 12 1:1.00 Group 1	Std 19 15360.00 0	Std 20 30720.00 0	Std 21 61440.00 0	Un 16 1:1.00 Group 1	Un 20 1:1.00 Group 1	Un 24 1:1.00 Group 1	Operator		+	×
Report	E	Std 05 5.000 Group 1	Std 05 5.000 Group 1	Std 05 5.000 Group 1	Neg 1 Group 1	Neg 1 Group 1	Pos 1 Group 1	Std 22 1.2E+5 Group 1	Std 23 2.5E+5 Group 1	Std 24 4.9E+5 Group 1	Neg 1 Group 1	Neg 1 Group 1	Pos 1 Group 1				~ /
	F	Std 06 6.000 Group 1	Std 06 6.000 Group 1	Std 06 6.000 Group 1	Pos 1 Group 1	Ctrl 01 Group 1	Ctrl 02 Group 1	Std 25 9.8E+5 Group 1	Std 26 2.0E+6 Group 1	Std 27 3.9E+6 Group 1	Pos 1 Group 1	Ctrl 10 Group 1	Ctrl 09 Group 1	Step by			1
	G	Std 07 7.000 Group 1	Std 07 7.000 Group 1	Std 07 7.000 Group 1	Ctrl 05 Group 1	Ctrl 04 Group 1	Ctrl 03 Group 1	Std 28 7.9E+6 Group 1	Std 29 1.6E+7 Group 1	Std 30 3.1E+7 Group 1	Ctrl 06 Group 1	Ctrl 07 Group 1	Ctrl 08 Group 1	Unit		nç	g/μL
	н	Std 08 8.000 Group 1	Std 08 8.000 Group 1	Std 08 8.000 Group 1	Blk 1 Group 1	Blk 1 Group 1		Std 31 6.3E+7 Group 1	Std 32 1.3E+8 Group 1		Blk 1 Group 1	Blk 1 Group 1		Sample group	S		\bigcirc
	Туре	Select			-		_			Provide a							
	0																
		Blank	Stan	dard	Qualit	y Ui	nknown	Posit	tive	Negative	e	Clear					
															L 11-1		

Fig 9

Note: The plate layout can be set up before or after the plate is read. For example, users can set all wells as "unknown", read the absorbance values, then set up the layout of the plate. The absorbance value data and layout are separate, users can create or modify a layout to analyze absorbance data that was captured in the past.

Note: the absorbance values of a test cannot be changed!

4.2 Parameter setting

After completing the plate layout settings, click "Run" on the left side bar (see Fig 10.)

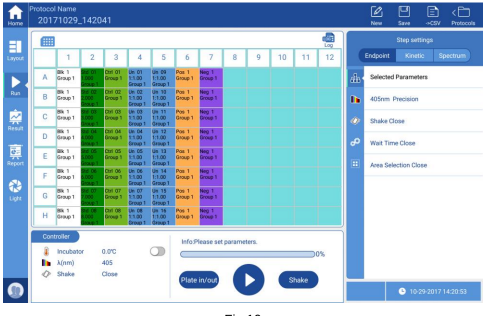


Fig 10

After clicking "Run" the lower window becomes a controller with the following functions:

- Incubator: The plate chamber can be set to a specified temperature, and the actual temperature is displayed.
- Wavelength: shows wavelength parameters of the current protocol.
- Shake: This function can be set to on or off.
- Progress bar: displays the running status of the current protocol.
- Plate in/out: control the motorized plate drawer to move it in and out; there is also a physical control button on the front of the instrument.
- Start/Stop: Used to start or stop a plate reading. There is also a physical button on the front of the enclosure for start/stop.
- Shake: This setting is independent from the shaking function in the settings menu, it is specifically used for plate defoaming.

In "Run" mode, the window on the right side allows choosing a specific measurement method: Endpoint, Kinetic or Spectrum, as well as reading parameters. See below table 2.

	Table 2	
Endpoint	Kinetic	Spectrum
Lucpont Selected Parameters Wavelength Lacktrian	Linetic L Selected Parameters L Wavelength LFast LPrecision L Wavelength LA1 (405) LA2 (450) LA2 (450) LA3 (492) LA4 (630) L Kinetic LTotal Time LTotal Time LTotal Time LNinetic region LNo. of readings LNumber	L Selected Parameters L Wavelength L Mode L Fast L Precision L Wavelength L Start Wavelength L End Wavelength L Step
└ Shake └Speed └Low └Medium └High └Type └Continuous └Pulsed └Time └ Wait Time at start └Area Selection	└Kinetic region └Shake └Speed └Low └Medium └High └Type └Continuous └Pulsed └Time └ Wait Time at start └ Area Selection	└ Shake └Speed └Low └Medium └High └Type └Continuous └Pulsed └Time └ Wait Time at start └Area Selection

For wavelength selection, Endpoint and Kinetic are configured with four wavelengths (defaults are 405nm, 450nm, 492nm and 630nm), users can click on and modify these wavelength values manually, but must be within the range of 200nm~1000nm. Spectral analysis allows setting a wavelength range within 200nm~1000nm.

A "Log" button appears at the top right corner of the plate layout window after the current protocol has been executed.

The "Log" is mainly used for recording the finishing time of each step of a protocol.

Note: Steps cannot be edited if a protocol has been executed, users can save it as a new one and then edit.

4.3 Reading a Plate

Click "Plate in/out" on the screen or press "Plate in/out" on the front of the enclosure to open the plate drawer, load an ELISA plate on the plate holder, paying attention to the orientation of the plate so the A1 well is in the correct location. See Fig 11.





Click button " P" on the display or press the "Start" button on the front of the instrument. If the protocol has already been run, a window will pop up to allow re-naming the protocol results file. Input a file name, click "ok" to run the protocol. The plate drawer will close, and during the reading process the screen will turn grey (see Fig 12) and all

buttons will become inoperable with the exception of the stop button "**U**". The user can also press the "Stop" button on the front of the housing to stop a run.

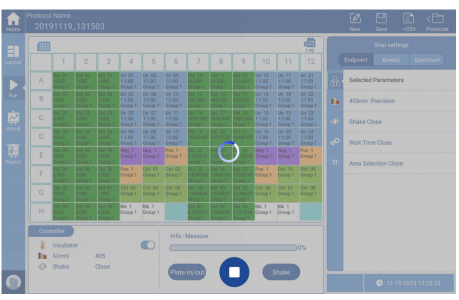


Fig 12

5. Result processing

The screen will stay in the "Run" interface after detecting samples and will display the original absorbance values measured. To view the analyzed results, switch to the "Result" interface by pressing the Result button on the side bar, see Fig 13.



Fig 13

Results are displayed according to the protocol settings and measurement method. The above Fig 13 shows results of an endpoint reading. Here, according to different types of detection modes, the Result is divided into three modes: Endpoint, Kinetic and Spectrum.

5.1 Endpoint result

As Fig 13 shows, data processing options for an endpoint result include: "Raw Data", "Blank Subtraction", "Basic Calculation", "Standard Curve", "Classification" and "Quality Control".

• Raw data: displays absorbance values of each well. If the plate was read using multiple wavelength, the user can switch to different wavelengths by pressing the

button " λ : 562" at the top, middle of the main interface. Blank Subtraction: An average blank absorbance value is obtained per the wells set for "blank" samples. This average value is then subtracted from the absorbance values for each sample well.

Note: 1. The Blank Subtraction button is only available if blanks have been set in the protocol layout interface.

2. When a protocol layout is set with blank samples, the absorbance values of "Basic Calculation", "Standard curve", "Classification", "Quality Control" and "Kinetic Analysis" all include the subtraction of the blank value.

- "Basic Calculation": the four basic functions of "+", "-", "x", "/" can be performed on the absorbance values at different wavelengths of the same well.
- "Standard Curve": Based on the concentrations set for the standard wells and the corresponding measured absorbance values, the instrument will generate a standard curve and use this for sample concentration calculations, see Fig 14.



Fig 14

If the curve fit is not closely matching the data points, the curve type can be changed in

the "Parameters" window on the right of the interface. Below are the 8 curve fitting types available:

- > Linear fitting (Minimum Standards required: 2)
- > Four Parameters Logistic (4PL) (Minimum Standards required: 4)
- Quadratic polynomial (<u>Minimum Standards required: 3</u>)
- > Cubic polynomial (Minimum Standards required: 4)
- > Quartic polynomial (Minimum Standards required: 5)
- > Point-to-point (Minimum Standards required: 2)
- Cubic Spline (<u>Minimum Standards required: 3</u>)
- Logit/Log (<u>Minimum Standards required: 2</u>)

The absorbance values and concentration values of the standards are set to a curve. The SmartReader supports the following types of curve fittings that can be chosen to best fit the standard curve. When R² value is closest to 1.0, this indicates an optimized curve fit.:

- > Linear/Linear (Linear fitting of absorbance value and corresponding concentration)
- Linear/Log (Linear fitting of absorbance value and the logarithm of the concentration)
- ➢ Log/Linear
- ➢ Log/Log
- Note: In order to generate a curve, standard wells must be set up on the Protocol Layout Interface. Some curve types require a minimum quantity of standards. In the event of a "curve fit failure", additional standards and concentration values may need to be included in the layout settings.
- Qualitative analysis: Samples can be qualitatively analyzed in relation to negative and positive references as set up in the plate layout. (see Fig 15). A formula is set up (on the right side of the interface), for calculating and marking positive "+", weak positive "t", or negative (no marking) sample wells automatically. Enter the appropriate values to create the formula for the "critical value", "weak value", and select if positive result will be > or < the critical value.

ne (91119_	_1315	03							÷			New	Save	->CSV	Protoco
		_									Print	Export	List		Parame	eters	
ut		1	2	3	4	5	6	7	8	9	10	11	12	Classification			
	А	Std 01 0.006 +	Std 01 -0.022 +	Std 01 -0.026 +	Un 01 1.350	Un 05 0.443 +	Un 06 0.036 +	Std 11 -0.039 +	Std 11 -0.045 +	Std 11 -0.040 +	Un 10 1.301	Un 14 0.413 +	Un 15 0.032 +	Critical Value	K ₁ :	×NC+K ₂ ×	PC+K ₃
	В	Std 02 0.275 +	Std 02 0.268 +	Std 02 0.273 +	Un 02 1.704	Un 05 0.442 +	Un 06 0.030 +	Std 12 0.251 +	Std 12 0.265 +	Std 12 0.253 +	Un 11 1.710	Un 14 0.422 +	Un 15 0.029 +	К,		1	1
	С	Std 04 0.529 +	Std 04 0.537 +	Std 04 0.521 +	Un 03 2.321	Un 05 0.432 +	Un 06 0.028 +	Std 14 0.530 +	Std 14 0.526 +	Std 14 0.524 +	Un 12 2.338	Un 14 0.430 +	Un 15 0.019 <mark>+</mark>				0
ılt	D		Std 05 0.954 +	Std 05 0.991 +	Un 04 2.677	Un 05 0.459 +	Un 06 0.028 +	Std 15 1.003	Std 15 0.994 L ⁺	Std 15 1.048	Un 13 2.741		Un 15 0.028 <mark>+</mark>				
rt	Е	Std 08 1.352	Std 08 1.345	Std 08 1.381	Neg 1 0.992 +	Neg 1 1.020 L ⁺	Pos 1 1.465	Std 18 1.399	Std 18 1.456	Std 18 1.464	Neg 2 0.979 L	Neg 2 0.929 +	Pos 2 1.663	K ₃			0
	F	Std 06 1.639	Std 06 1.711	Std 06 1.743	Pos 1 1.362	Ctrl 01 0.534 +	Ctrl 01 0.582 +	Std 16 1.780	Std 16 1.713	Std 16 1.795	Pos 2 1.723	Ctrl 05 0.533 +	Ctrl 05 0.511 +	Weakly Value	±k	<₄%×Criti	cal Valu
	G	Std 09 2.251	Std 09 2.289	Std 09 2.326	Ctrl 02 0.297 +	Ctrl 03 2.312	Ctrl 04 0.468 +	Std 17 2.764	Std 17 2.759	Std 17 2.752	Ctrl 09 0.274 +	Ctrl 10 2.323	Ctrl 11 0.067 +	K4			1
	н	Std 07 2.596	Std 07 2.700	Std 07 2.682	Blk 1 -0.009 <mark>+</mark>	Blk 1 0.009 +		Std 19 2.314	Std 19 2.301	Std 19 2.298	Bik 2 0.020 🕇	Blk 2 -0.020 +		Positive	> (Crit	tical Valu
	Calcu	lation										1		+ Positive	L+	Low Pos	sitive
	ource			B					6								
	Raw	Data		6 Blank		Basic		Standard		***		✓ Quality					
				Subtractior		Calculation		Curve		lassificatio		Control		~	B 09	31-2009 1	2-1 5-20

Fig 15

Note: The Protocol layout interface must be set with at least one positive or negative well, or the "Classification" button will not be available. See Fig. 15

 Quality Control: Click "Quality Control" button, (see Fig 16), the main display area will turn out to be a list, it displays status of the quality control wells which have set previously, the instrument will mark the samples according to conditions set in "Parameters "Parameters".

			Prin	nt Export List		Param	eters	
No.	Sample	Input	Value	Result	Quality Contro	bl		
1	Ctrl 01	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Passed	Input			
2	Ctrl 02	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Passed				
3	Ctrl 10	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Failed	Target			1.0
4	Ctrl 09	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Failed	SD			0.5
5	Ctrl 05	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Passed				
6	Ctrl 04	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Passed			Ad	d Ru
7	Ctrl 03	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Passed				
8	Ctrl 06	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Passed				
9	Ctrl 07	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Passed				
10	Ctrl 08	Target:1.000 SD:0.500	Cofficient0.5 Upper2.0 Lower0.0	Failed				
Calcul		Blank Basic Calculation	Standard Curve Classification	Quality Control		• • • •	-31-2009 1	10.17

Fig 16

Note: protocol layout interface must be set with "Quality" wells, otherwise, this button will be not available.

5.2 Kinetics

After a kinetic type assay, the results screen will display small graphs inside the squares instead of absorbance values (see Fig 17).

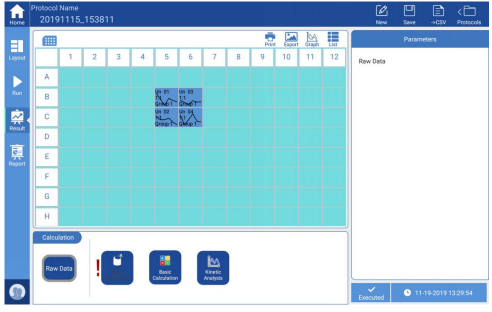


Fig 17

To see the detailed absorbance results for a specific well, select the target well, then click the "Graph" button at the top right to enlarge the kinetic curve (see Fig 18). Click "Back" to return to the main interface.



Fig 18

The "Calculation" window at the bottom offers four selections: "Raw Data", "Blank Subtraction", "Basic Calculation" and "Kinetic Analysis" (see Fig 18).

For functions and algorithms of "Raw Data", "Blank Subtraction" and "Basic Calculation",

please see Section 5.1.

Clicking the "Kinetic Analysis" button, the window on the right side will offer different calculation options, (see Fig 19).

The calculations available include:

- > Average/SD/CV%
- Integral
- Baseline Subtraction
- Select Single Reading
- Select Reading Range
- Maximum Rate
- Maximum (Peak)

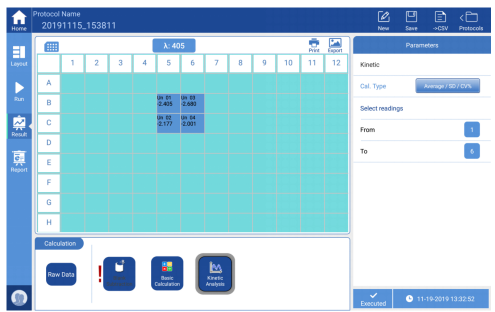


Fig 19



5.3 Spectral Analysis

See Fig 20 (above), select the target wells click "Graph" button to enlarge the spectral curve, the absorbance of each wavelength shown on the curve. Users can click the "Back" button to return to the main interface.

Click "Spectral Analysis", see Fig 21, for further analysis.

Calculations available:

- Spectral Maximum: read the maximum value greater than the threshold value within a set range.
- Spectral Normalization: set the spectral range, take the maximum absorption peak as 1, the remaining values will be converted into percentage based on this criterion.
- > Ratio within the Spectrum: set two wavelength values $\lambda 1$ and $\lambda 2$, calculate the value of $\lambda 1/\lambda 2$.
- Select Wavelength Range: read measurement values within a set wavelength range.

Select Single Wavelength : read the measurement value of a single wavelength.

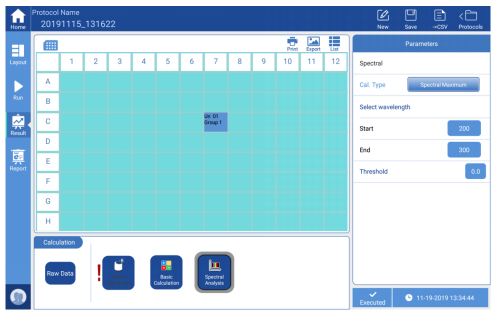


Fig 21a

	T		roplate						Prin	t Export	Graph	List			Paramet	ers	
4	1	2	3	4	5	6	7	8	9	10	11	12	Raw [Data			
А	BIK 1 Group	BIK DV															
В	Bik 1 Group 1	BIK 1 GNO P															
С	Blk M Group 1	Bik 1 Gloup															
D	Bik 1 Graup																
E	Bik W Group	Bik TA															
F		•															
G																	
Н																	



Description of each button on the upper right corner of the main display area:



Print the current content of the main display area to a connected thermal printer



Export the current content of the main display area as an image file to a flash drive.



Choose a well to view the kinetic curve, the "Graph" button can enlarge the curve, see Fig 17, "Back" is to return to the main interface.

Switch from plate layout view to a data list display, click the "Plate" button in the upper right corner to switch back to the plate layout display.



Restore the zoom function



Return to the previous interface

6. Report exporting

Both processed data and raw data can be exported, click the "Report" button on the left side menu, see Fig 22.

f Home	Protoco 2019		_1315	03											New	Contraction Save	.⇒csv	< 🛅 Protocols
						λ: 40	05		_				List			Output C	ontent	
Layout		1	2	3	4	5	6	7	8	9	10	11	12	Info				
	А	Std 01 0.186	Std 01 0.156	Std 01 0.152	Un 01 1.529	Un 05 0.623	Un 06 0.216	Std 11 0.153	Std 11 0.147	Std 11 0.152	Un 10 1.495	Un 14 0.606	Un 15 0.225	Ge	neral int	ormation		~
Run	В	Std 02 0.454	Std 02 0.447	Std 02 0.453	Un 02 1.884	Un 05 0.621	Un 06 0.210	Std 12 0.444	Std 12 0.458	Std 12 0.446	Un 11 1.903	Un 14 0.616	Un 15 0.222	Ins	strumen	tinformati	on	~
2	С	Std 04 0.709	Std 04 0.717	Std 04 0.700	Un 03 2.500	Un 05 0.612	Un 06 0.208	Std 14 0.724	Std 14 0.719	Std 14 0.717	Un 12 2.531	Un 14 0.624	Un 15 0.212	_				
Result	D	Std 05 1.152	Std 05 1.134	Std 05 1.170	Un 04 2.856	Un 05 0.639	Un 06 0.208	Std 15 1.197	Std 15 1.187	Std 15 1.241	Un 13 2.934	Un 14 0.629	Un 15 0.221	Pro	otocol p	arameters		~
Report	Е	Std 08 1.531	Std 08 1.524	Std 08 1.560	Neg 1 1.171	Neg 1 1.199	Pos 1 1.644	Std 18 1.592	Std 18 1.650	Std 18 1.657	Neg 2 1.172	Neg 2 1.123	Pos 2 1.856	Layou	ut defini	tions		~
кероп	F	Std 06 1.819	Std 06 1.891	Std 06 1.922	Pos 1 1.541	Ctrl 01 0.714	Ctrl 01 0.761	Std 16 1.973	Std 16 1.907	Std 16 1.988	Pos 2 1.916	Ctrl 05 0.726	Ctrl 05 0.705	RawD	Data			~
	G	Std 09 2.430	Std 09 2.468	Std 09 2.505	Ctrl 02 0.477	Ctrl 03 2.492	Ctrl 04 0.648	Std 17 2.957	Std 17 2.952	Std 17 2.945	Ctrl 09 0.454	Ctrl 10 2.503	Ctrl 11 0.246	Resul	It			
	н	Std 07 2.776	Std 07 2.879	Std 07 2.861	Blk 1 0.170	Blk 1 0.188		Std 19 2.507	Std 19 2.495	Std 19 2.491	Blk 2 0.213	Blk 2 0.172		Bla	ank Subt	raction		~
	File	Туре												Ba	sic Calc	ulation		~
		x		c 🗏		PE		ΓΞ								Outp	ut	
0		XL	5	CSV		PDF		тхт		ļ		Print		Execu		G 11-	19-2019 1	3:36:11

Fig 22

Choose the desired format in the "File Type" window at the bottom, four formats are available:

- XLS
- CSV
- PDF
- TXT

Choose the content to export in "Output Content" on the right side, " $\sqrt{}$ " will appear next to the selected content, then click "Output" to export data to a flash-drive.

"Print": The print function can only be used to print the instrument's basic information, including instrument serial number, software version etc.

7. Power off

Press the "Plate in/out" button to remove the ELISA plate from the Reader, then press the button again to insert the empty plate holder back into the chamber.

Turn off the power switch on the back of the instrument.

Chapter 6 Maintenance, storage, transportation

1. Maintenance

- Keep the installation environment dry and clean to prevent moisture, corrosion, and strong electromagnetic interference sources.
- The SmartReader Instrument is calibrated before it leaves the factory. There are no user-serviceable parts inside, so please do not attempt to open the housing. If any defects or operation problems arise, please contact your distributor or Accuris Instruments at 908-769-5555 or info@accuris-usa.com.
- Do not switch on and off the power abruptly, as this can damage the electronics.
- Make sure to connect the SmartReader to an appropriate power outlet (voltage and Hz) as stated on the instrument rating plate.
- Maintenance list

Content	Daily	Weekly	Annually	When needed
Make sure the instrument powers off correctly				\checkmark
Keep the instrument away from dust	\checkmark			
Remove spilled solutions right away to prevent damage, then clean it with distilled water.	\checkmark			
If any surface has been contaminated with				
biohazardous materials, sterilize it using a	\checkmark			
disinfectant.				
Clean instrument housing regularly.		\checkmark		
Clean the plate holder when necessary.				
Maintenance				\checkmark

2. Storage and transportation

- When not in use, store the instrument at a temperature range -10°C~40°C, relative humidity less than 80%, without corrosive gas and with good ventilation.
- Prevent from heavy vibrations and humidity during transportation.

No	Trouble description	Possible reason	Solution	
1	The Microplate Reader cannot be started	Power supply failure	 a. Check that there is power to the instrument b. Make sure that all power cords are connected properly c. Confirm that the wall voltage is correct. 	
2	"Communication timeout" during self-checking	Instrument not working	Restart the instrument and try again; if the problem persists, please contact your distributor or Accuris Instruments.	
3	"E913, E923, E933, E943" during self-checking	Insufficient light intensity	Please contact your distributor or Accuris Instruments.	
4	"E912, E922, E932, E942" when self-checking	Light intensity is too strong	Please contact your distributor or Accuris Instruments.	
5	"E911, E921, E931, E941" when self-checking	Excessive dark current	Please contact your distributor or Accuris Instruments.	
6	"E612, E622, E632, E642" when self-checking	Detection module failure	Please contact your distributor or Accuris Instruments.	
7	"E401, E403, E415, E425, E435, E445" when self-checking	Motor failure	Please contact your distributor or Accuris Instruments.	
8	"E011~E056" when self-checking	Incubation failure	Please contact your distributor or Accuris Instruments.	
9	Test results are greatly deviated or all are zero	Xenon lamp damaged	Restart the instrument and try again; if the problem persists, please contact your distributor or Accuris Instruments.	
10	Elisa plate holder not properly moving in or out	Something is obstructing the movement	Check for any objects around the plate holder, if the plate cover is raised, or if the plate is not inserted properly	
11	Abnormal noise occurs during running	The Elisa plate is not correctly seated or the plate cover is not seated	 a. Check Elisa plate b. If the noise still occurs there when running without a plate, restart the instrument c. If noise persists, please contact your distributor or Accuris Instruments. 	

Chapter 7 Trouble shooting

12	Test results are unstable	Light path failure	Check that the plate is properly seated, if liquid has spilled out and whether the front door is working correctly, then re-start the instrument. If problem still occurs, contact your distributor or Accuris Instruments
13	Instrument stops running during detection	Communication breakpoint	Press "stop", restart the detection

No.	ltem	Туре	Unit	Qty	Remarks
1	Power cord		PCS	1	
2	AC/DC Power Adapter	24V 160W	PCS	1	
3	Mouse	Logitech	PCS	1	
4	Performance test statement		PCS	1	
5	Operation Manual		PCS	1	
6	Packing List		PCS	1	
7	Flash Drive		PCS	1	
8	Certification		PCS	1	
9					

Chapter 8 Accessories