Part 1 of 6	Bio/Data Corp.	Chrono-log Corp.	Chrono-log Corp.
	Mary Brown customer.service@biodatacorp.com Horsham, PA	Sal Pema sal@chronolog.com Havertown, PA 610, 952, 1130, or 900, 347, 6665, www.obronolog.com	Sal Pema sal@chronolog.com Havertown, PA 610, 952, 1120, or 900, 247, 6665, www.ehronolog.com
Instrument name/First year cold	215-441-4000 or 800-257-3282 www.biodatacorp.com	610-853-1130 or 800-247-6665 www.chronolog.com	610-853-1130 or 800-247-6665 www.chronolog.com
Instrument name/First year sold List price/Model type	PAP-8E Platelet Aggregometer/2005	Optical Aggregation Systems Models 490 4+, 490 4+4/2017 \$11,268-\$26,795/benchtop	Whole Blood Optical Lumi-Aggregation System 700-2, 700-4/2006 \$19,909-\$42,100/benchtop
District Moder type Dimensions (H × W × D)/Weight/Instrument footprint No. of units in clinical use in U.S./Outside U.S. (countries) Composition of installs: Hospital lab/Reference lab/Other Targeted daily, monthly, annual test volume	\$22,990/benchtop 21.5-25.5 × 19.5 × 21.7 in./40 lbs./4 sq. ft. >500/>300 (worldwide) 80%/15%/5% (unspecified)	per each 4-channel module: 8.5 × 14 × 15 in./19.3 lbs./1.5 sq. ft. —	per each 2-channel module: 8.5 × 14 × 18 in./40 lbs./1.75 sq. ft. — —
Operational type	batch, random access	discrete	discrete
Country where analyzer designed/Manufactured Company manufactures instrument	U.S./U.S. yes (also sold by möLab, Sentinel Diagnostics, Werfen, Alpha Labs, Sysmex, Analis)	U.S./U.S. yes (also sold via distribution partners)	U.S./U.S. yes (also sold via distribution partners)
FDA-approved clotting-based tests FDA-approved chromogenic tests	=		_
FDA-approved immunologic tests Other FDA-approved tests	ristocetin cofactor activity, HIT/HIPA, RIPA, others routine platelet aggregation testing, platelet activation, spontaneous platelet aggregation, sticky platelets, washed platelets, others	platelet agglutination with ristocetin LTA platelet aggregation, ristocetin cofactor assay	platelet agglutination with ristocetin whole blood impedance platelet aggregation, LTA platelet aggregation, platelet dense granule ATP release, ristocetin cofactor assay
User-defined tests in clinical use	<u>-</u>	LTA platelet aggregation, ristocetin cofactor assay	whole blood impedance platelet aggregation, LTA platelet aggregation, platelet dense granule secretion, ristocetin cofactor assay
Tests in development or awaiting FDA 510(k) clearance	_	_	
Methodologies supported	turbidimetric, immunologic (agglutination, Ab-Ag tests)	turbidimetric for measuring platelet aggregation in PRP and ristocetin cofactor assay	turbidimetric for measuring platelet aggregation in PRP and ristocetin cofactor assay, impedance for measuring platelet aggregation in whole blood
Number of different measured assays onboard simultaneously Number of different assays programmed and calib. at one time		4–8 4–8	2–4 2–4
No. of user-definable (open) channels/No. active simultaneously Factor assays require manual manipulation or dilutions	8/8 yes (manual manipulation and dilutions)	4–8/4–8 yes (manual dilutions)	2–4/2–4 yes (manual dilutions)
Test throughput per hour/Assay run time	80 (up to 720 samples in throughput)/6 min. average	6 (24–48 tests in throughput)/5 min. minimum	6 (12–24 tests in throughput)/6 min. minimum
Design of sample-handling system Operates on whole blood or spun plasma	manual spun plasma	manual spun plasma	manual whole blood and spun plasma
Reagent type Reagent barcode-reading capability	open reagent system (liquid, lyophilized [reconstituted manually]) yes, for all tests	self-contained multiuse vials; open reagent system (liquid, lyophilized [reconstituted manually]) no	self-contained multiuse vials; open reagent system (liquid, lyophilized [reconstituted manually]) no
No. of reagent containers held onboard/Reagents ready to use Reagent lot tracking/Reagent inventory/Reagents refrigerated onboard	none/requires operator prehandling yes/no/no	—/requires operator prehandling no/no/no	/requires operator prehandling no/no/no
Reagents, consumables loaded without interrupting testing Instrument uses proprietary or third-party reagents Maximum time same lot number of reagents can be used	yes (reagents and consumables) user's option 2 years	yes (consumables) proprietary reagents 18 months–3 years	yes (consumables) proprietary reagents 18 months–3 years
Walkaway capability/Walkaway duration	yes/8 specimens or 8 tests	yes/5 min. or 4–8 specimens or 4–8 tests	yes/6 min. or 2–4 specimens or 2–4 tests
Min.—max. specimen volume that can be aspirated at one time Min. sample volume required for PT/PTT/Factor VIII activity Types of disposables used	— — — micro test tubes, micro stir bars, pipette tips, sample tubes,	250–500 µL cuvettes, stir bars, pipette tips	225–500 µL cuvettes, stir bars, disposable electrodes, pipette tips
	sample tube caps		
Primary tube sampling supported/Pierces caps on primary tubes Accommodates most standard tube sizes/Nonstandard sizes	no/no no/no	no/no no/no	no/no no/no
Sample barcode-reading capability/Autodiscrimination Auto tracks product volume/Measures number of tests remaining	no/yes	no/no	no/no
Short sample detection	no/yes no	no/no no	no/no no
Clot detection as preanalytical variable in plasma sample Auto detects adequate reagents for aspiration or analysis	no no	no no	no no
Detection or quantitation for hemolysis, turbidity, icterus, lipemia	detection for hemolysis, turbidity, icterus, lipemia	no	no
Dilutes patient samples onboard Automatic rerun capability/Auto reflex testing capability	no no/no	no yes/no	no yes/no
Lag time during which hypercoagulable sample not detected	no	no	no
User can adjust reagent volumes/Sample volumes User can adjust No. of reagents/Sources of reagents	yes/yes yes/yes	yes/yes yes/yes	yes/yes yes/yes
User can adjust incubation times/Reading times	yes/yes	yes/yes	yes/yes
Read time extended for prolonged clotting times Autocalibration/Calibrants stored onboard	no yes/no	yes (selectable on menus) yes/no	yes (selectable on menus) yes/no
Multipoint calibration supported/Recommended frequency	yes/annually	yes/annually	yes/annually
Stat time to complete all analytes/Throughput per hour for: • PT alone	_	_	_
• PT, PTT	_	_	_
Fibrinogen Factor VIII activity assay	_		_
• D-dimer	_	_	_
Time delay from ordering stat to aspiration of sample How labs get LOINC codes for results	— email query	email query	— email query
Onboard real-time QC/Onboard software capability to review QC Information that can be barcode-scanned on instrument	no/yes reagent lot No.	yes/no barcode scanning not offered	yes/no barcode scanning not offered
Compatible with laboratory automation systems	no	no	no
Data-management capability/LIS or EHR systems interfaced Interface supplied by instrument vendor	onboard/none no	onboard/none no	onboard/none no
Results transferred to LIS as soon as test time complete	no	no	no
Bidirectional interface capability	no	no	no
Remote servicing provided/UPS backup power supply Instrument connections to transfer information Interface standards supported	no/no 	no/no 	no/no — —
Information transferred to data-management software	T	-	-
Avg. time for basic user training Approximate scheduled maintenance time	1 day (at customer site or virtually) weekly: 15 minutes; monthly: 30 minutes	— preventive maintenance and calibration by clinical engineering recommended annually	1.5 days (at customer site) preventive maintenance and calibration by clinical engineering recommended annually
Maintenance records kept onboard	NO (\$2.050	yes	yes
Warranty with purchase/Annual service contract cost (24/7)	yes/\$2,050	yes/— (cost dependent on contract)	yes/— (cost dependent on contract)
Distinguishing features (supplied by company)	eight independently operated channels, nine results reported per channel, standard micro volumes; preloaded routine and special test parameters, user-defined test templates and protocols, two- year warranty; includes all-in-one wireless computer, PAP-8E		3 instruments in 1: whole blood/impedance and PRP/LTA aggregometer plus luminometer to measure platelet ATP release; continuously monitors and regulates temperature and stirring; optical calibration can be performed by laboratory
Note: a dash in lieu of an answer means company did not answer question or question is not applicable	year warranty; includes all-in-one wireless computer, PAP-8E system software, electronic pipette, and Microsoft Office Suite		personnel using no-cost water samples

Part 2 of 6	Diagnostica Stago Nichole Howard nichole.howard@us.stago.com Parsippany, NJ 800-222-2624 www.stago-us.com	Diagnostica Stago Nichole Howard nichole.howard@us.stago.com Parsippany, NJ 800-222-2624 www.stago-us.com	Diagnostica Stago Nichole Howard nichole.howard@us.stago.com Parsippany, NJ 800-222-2624 www.stago-us.com
Instrument name/First year sold	STA Compact Max 3/2023	STA Satellite/2010	STA-R Max 3/2023
List price/Model type Dimensions (H × W × D)/Weight/Instrument footprint No. of units in clinical use in U.S./Outside U.S. (countries) Composition of installs: Hospital lab/Reference lab/Other	\$164,794/benchtop 27.75 × 38.18 × 28.73 in./309 lbs./7 sq. ft. ~1,850/~4,100 (France, Spain, UK, Germany, Denmark, others) ~96%/3%/1% (veterinary labs, pharmaceutical companies,	\$58,935/benchtop 27.4 × 21.1 × 25.5 in./72 lbs./4 sq. ft. ~550/~2,100 (France, Spain, UK, Germany, Denmark, others) ~98%/0/2% (veterinary labs, academic research)	\$250,770/floor standing 49.2 × 50.3 × 32.2 in./564 lbs./26.8 sq. ft. ~400/~1,800 (France, Spain, UK, Germany, Denmark, others) ~90%/7%/3% (veterinary labs, pharmaceutical companies,
Targeted daily, monthly, annual test volume	academic research/educational labs) daily: >30 (moderate-volume laboratories); monthly: 900; annual: >5,000	daily: ~40; monthly: <900; annual: <13,000	academic research/educational labs) daily: >100 (moderate- to high-volume laboratories); monthly: >2,500; annual: >25,000
Operational type Country where analyzer designed/Manufactured Company manufactures instrument	continuous random access France/France yes	random access France/France yes	continuous random access France/France yes
FDA-approved clotting-based tests	PT, APTT, TT, fibrinogen, reptilase, factors, proteins C and S,	PT, APTT, fibrinogen	PT, APTT, TT, fibrinogen, reptilase, factors, proteins C and S,
FDA-approved chromogenic tests	lupus anticoagulant, DRVVT (screen and confirm) anti-FXa (UFH and LMWH), antithrombin, protein C, plasminogen, FVIII chromogenic	anti-FXa (UFH and LMWH), antithrombin	lupus anticoagulant, DRVVT (screen and confirm) anti-FXa (UFH and LMWH), antithrombin, protein C, plasminogen, FVIII chromogenic
FDA-approved immunologic tests	D-dimer, antithrombin antigen, free protein S, total protein S, wF antigen (all microlatex)	D-dimer	D-dimer, antithrombin antigen, free protein S, total protein S, vWF antigen (all microlatex) STA NeoPTimal protime reagent with ISI ~1.0
Other FDA-approved tests User-defined tests in clinical use	STA NeoPTimal protime reagent with ISI ~1.0 APCR, other clotting, chromogenic, and immunological tests with user-defined applications	_	APCR, other clotting, chromogenic, and immunological tests with user-defined applications
Tests in development or awaiting FDA 510(k) clearance	-	-	-
Methodologies supported Number of different measured assays onboard simultaneously	mechanical clot detection, chromogenic, immunologic (microlatex) 80	mechanical clot detection, chromogenic, immunologic (microlatex) 80	mechanical clot detection, chromogenic, immunologic (microlatex) 200
Number of different assays programmed and calib. at one time No. of user-definable (open) channels/No. active simultaneously Factor assays require manual manipulation or dilutions		80 80/80	200 200/200
Test throughput per hour/Assay run time	110 (2 tests in throughput)/4.6–7.3 min. (avg. 4.7 min.)	36 (3 tests in throughput for PT, APTT, fibrinogen)/4.6–7.3 min. (avg. 4.7 min.)	280 (2 tests in throughput)/4.6–7.3 min. (avg. 4.7 min.)
Design of sample-handling system Operates on whole blood or spun plasma	continuous loading sample drawer with continuous random access spun plasma	continuous sample loading with positive sample identification, removable sample carousel for 20 primary tubes spun plasma	rack with continuous specimen access spun plasma
Reagent type	self-contained single-use and multiuse vials; open reagent system (liquid, lyophilized [reconstituted manually])	self-contained single-use and multiuse vials; open reagent system (liquid, lyophilized [reconstituted manually])	self-contained single-use and multiuse vials; open reagent system (liquid, lyophilized [reconstituted manually])
Reagent barcode-reading capability No. of reagent containers held onboard/Reagents ready to use Reagent lot tracking/Reagent inventory/Reagents refrigerated onboard	yes, for all tests 45/variable (reagent specific) yes/yes (15°-19°C)	yes, for all tests 16/variable (reagent specific) yes/yes (15°-19°C)	yes, for all tests 75/variable (reagent specific) yes/yes/yes (15°-19°C)
Reagents, consumables loaded without interrupting testing Instrument uses proprietary or third-party reagents Maximum time same lot number of reagents can be used	yes (consumables) user's option (same capabilities when third-party reagents used) 18 months	yes (consumables) user's option (same capabilities when third-party reagents used) 18 months	yes (consumables) user's option (same capabilities when third-party reagents used) 18 months
Walkaway capability/Walkaway duration	yes/96 specimens or 12 tests	yes/20 specimens or 12 tests	yes/215 specimens or 32 tests
Minmax. specimen volume that can be aspirated at one time Min. sample volume required for PT/PTT/Factor VIII activity	5–100 μL 50 μL/50 μL/50 μL	5–100 μL 50 μL/50 μL/50 μL	5–100 μL 50 μL/50 μL/50 μL
Types of disposables used	cuvettes, stir bars, cleaner solution	cuvettes, stir bars, cleaner solution	cuvettes, stir bars, cleaner solution
Primary tube sampling supported/Pierces caps on primary tubes Accommodates most standard tube sizes/Nonstandard sizes	yes/yes yes/no	yes/no yes/no	yes/yes yes/no
Sample barcode-reading capability/Autodiscrimination Auto tracks product volume/Measures number of tests remaining	yes (Interleaved 2 of 5, UPC, Codabar)/— yes/no	yes (Interleaved 2 of 5, UPC, Codabar)/no yes/no	yes (Interleaved 2 of 5, UPC, Codabar)/— yes/no
Short sample detection Clot detection as preanalytical variable in plasma sample	yes no	yes no	yes no
Auto detects adequate reagents for aspiration or analysis Detection or quantitation for hemolysis, turbidity, icterus, lipemia Dilutes patient samples onboard	yes (aspiration and analysis) yes	yes (aspiration and analysis)	yes (aspiration and analysis) yes
Automatic rerun capability/Auto reflex testing capability	yes yes/yes	yes yes/yes	yes yes/yes
Lag time during which hypercoagulable sample not detected	no vec/vec	no voc/voc	no weekee
User can adjust reagent volumes/Sample volumes User can adjust No. of reagents/Sources of reagents	yes/yes yes/yes	yes/yes yes/yes	yes/yes yes/yes
User can adjust incubation times/Reading times	yes/yes	yes/yes	yes/yes
Read time extended for prolonged clotting times Autocalibration/Calibrants stored onboard	yes (selectable on menus) yes/yes	yes (selectable on menus) yes/yes	yes (selectable on menus) yes/yes
Multipoint calibration supported/Recommended frequency	yes/6 months	yes/6 months	yes/6 months
Stat time to complete all analytes/Throughput per hour for: • PT alone • PT, PTT	<6 minutes/~150 specimens <6 minutes/~100 specimens	<6 minutes/~50 specimens <6 minutes/~40 specimens	<6 minutes/~320 specimens <6 minutes/~280 specimens
• Fibrinogen	<6 minutes/~100 specimens	<6 minutes/~40 specimens	<6 minutes/~180 specimens
Factor VIII activity assay D-dimer	<6 minutes/~60 specimens 7 minutes/~60 specimens	T minutes / 6 ansoimans	<6 minutes/~180 specimens 7 minutes/~150 specimens
Time delay from ordering stat to aspiration of sample	<15 seconds	7 minutes/~6 specimens <15 seconds	<15 seconds
How labs get LOINC codes for results Onboard real-time QC/Onboard software capability to review QC	email query, LOINC codes available in STA Coag Expert yes/yes	email query yes/yes	email query, LOINC codes available in STA Coag Expert yes/yes
Information that can be barcode-scanned on instrument	specimen identifier, reagent lot No., quality control ranges, calibrator values	specimen identifier, reagent lot No., quality control ranges, calibrator values	specimen identifier, reagent lot No., quality control ranges, calibrator values
Compatible with laboratory automation systems Data-management capability/LIS or EHR systems interfaced	no onboard/Cerner, Meditech, Clinisys, SCC, McKesson, Epic	no onboard/Cerner, Meditech, Clinisys, SCC, McKesson, Epic	yes (Stago, Abbott, Beckman Coulter, Cerner, Inpeco, QuidelOrtho, Roche, Siemens) onboard/Cerner, Meditech, Clinisys, SCC, McKesson, Epic
Interface supplied by instrument vendor Results transferred to LIS as soon as test time complete Bidirectional interface capability	contract dependent yes yes (broadcast download and host query)	contract dependent yes yes (host query)	contract dependent yes yes (broadcast download and host query)
Remote servicing provided/UPS backup power supply Instrument connections to transfer information	yes/yes data-management system, which in turn connects to LIS; directly to LIS; directly to lab automation system	no/yes directly to LIS	yes/yes data-management system, which in turn connects to LIS; directly to LIS; directly to lab automation system
Interface standards supported Information transferred to data-management software Avg. time for basic user training Approximate scheduled maintenance time Maintenance records kept onboard	ASTM 1394-91, ASTM 1381 device unique identifier, patient ID, specimen ID, result, QC identifier 3.5 days (at vendor office) daily: 10 minutes; weekly: 10 minutes; monthly: none yes	ASTM 1394-91, ASTM 1381 device unique identifier, patient ID, specimen ID, result, QC identifier 2.5 days (at vendor office) weekly: <15 minutes; monthly: <15 minutes no	ASTM 1394-91, ASTM 1381 device unique identifier, patient ID, specimen ID, result, QC identifier 3.5 days (at vendor office) daily: 10 minutes; weekly: 10 minutes; monthly: none yes
Warranty with purchase/Annual service contract cost (24/7)	yes/— (cost dependent on contract)	yes/— (cost dependent on contract)	yes/— (cost dependent on contract)
Distinguishing features (supplied by company)	viscosity-based, mechanical clot detection; precalibrated D-dimer and fibrinogen reagents, full complement of lupus anticoagulant testing; STA Coag Expert and Coag. One data managers deliver full	viscosity-based, mechanical clot detection; precalibrated D-dimer and fibrinogen reagents, 2-mL 24-hour quality control for PT, APTT, fibrinogen; small footprint for low-	viscosity-based, mechanical clot detection; precalibrated D-dimer and fibrinogen reagents, full complement of lupus anticoagulant testing; STA Coag Expert and Coag. One data managers deliver full
Note: a dash in lieu of an answer means company did not answer question or question is not applicable	autoverification, repeat/reflex testing, auto upload of QC to peer group and of CPR test counts	throughput labs with limited space	autoverification, repeat/reflex testing, auto upload of QC to peer group and of CPR test counts

Part 3 of 6	HemoSonics LLC Jeff Light _jlight@hemosonics.com Durham, NC	LABiTec LAbor BioMedical Technologies GmbH M. Schramm info@labitec.de Ahrensburg, Germany	LABiTec LAbor BioMedical Technologies GmbH M. Schramm info@labitec.de Ahrensburg, Germany
	401-741-3265 www.hemosonics.com	011-49-4102-47950 www.labitec.com	011-49-4102-47950 www.labitec.com
Instrument name/First year sold	Quantra Hemostasis Analyzer/2019	CoaDATA 2004 and 4004/—	CoaLAB 1000/—
List price/Model type	—/portable, benchtop	—/benchtop	—/benchtop
Dimensions (H × W × D)/Weight/Instrument footprint No. of units in clinical use in U.S./Outside U.S. (countries)	19 × 14 × 12 in./36 lbs./1 sq. ft. >100/>100 (Germany, France, England, Spain, Austria, Japan, Hong Kong, Australia, Italy, Portugal, Belgium)	10 × 13 × 3.5 in./8.6 lbs./0.92 sq. ft. —/>1,500 (worldwide [except U.S., Canada])	19.6 × 30.7 × 23.6 in./70.5 lbs./5 sq. ft. —/— (worldwide [except U.S., Canada])
Composition of installs: Hospital lab/Reference lab/Other Targeted daily, monthly, annual test volume Operational type	10%/0/90% (point of care [operating room, ICU, stat lab, more]) daily: 15–20; monthly: 1–200; annual: 1,200–2,400 random access, continuous random access	— batch	
Country where analyzer designed/Manufactured	U.S./U.S.	Germany/Germany	Germany/Germany
Company manufactures instrument	yes (also sold via distribution partners in parts of Europe, Asia)	yes (also sold via OEM distribution, local distributors)	yes (also sold via OEM distribution, local distributors)
FDA-approved clotting-based tests	QPlus cartridge: clot time, clot time with heparinase, clot time ratio, clot stiffness, more; QStat cartridge: clot time, clot stability to lysis, clot stiffness, platelet contribution to clot stiffness, more	_	_
FDA approved chromogenic tests	_	_	_
FDA-approved immunologic tests Other FDA-approved tests	_	_	_
User-defined tests in clinical use	_	_	_
Tests in development or awaiting FDA 510(k) clearance			
Methodologies supported Number of different measured assays onboard simultaneously	clot detection, sonic estimation of elasticity via resonance (SEER) sonorheometry 11	clot detection, mechanical and optical; photometric with mechanical stirring, turbodensitometric; chromogenic; immunologic (photometric) 15	clot detection, mechanical and optical; photometric with mechanical stirring, turbodensitometric; chromogenic; immunologic (photometric) 15
Number of different assays programmed and calib. at one time	no calibration required	15	50
No. of user-definable (open) channels/No. active simultaneously Factor assays require manual manipulation or dilutions	_	yes (manual manipulation and dilutions)	50/15 —
Test throughput per hour/Assay run time	5–6 (25–36 tests in throughput)/7–60 min. (avg. 12.5 min.)	CoaDATA 2004: ~60 PT tests (1 test in throughput); CoaDATA 4004: ~120 PT tests (1 test in throughput)/—	120 PT tests/—
Design of sample-handling system	sealed room-temperature-stable cartridges accept a standard 3.2% citrate blue top tube, manually affixed to cartridge	semiautomated analyzer with 2 and 4 channels	cuvette ring, sample cups
Operates on whole blood or spun plasma Reagent type	whole blood self-contained single-use cartridges (lyophilized [reconstituted	spun plasma self-contained single-use vials; open reagent system (liquid,	spun plasma self-contained single-use vials; open reagent system (liquid,
Reagent barcode-reading capability	manually]) yes, for all tests	lyophilized [reconstituted manually]) yes, for all tests	lyophilized [reconstituted manually]) yes, for some tests
No. of reagent containers held onboard/Reagents ready to use Reagent lot tracking/Reagent inventory/Reagents refrigerated onboard	1 cartridge/yes yes/no/no (20°-25°C)	4/variable (reagent specific) yes/—/no	15/yes yes/yes/no
Reagents, consumables loaded without interrupting testing	no	yes (reagents and consumables)	yes (reagents)
Instrument uses proprietary or third-party reagents Maximum time same lot number of reagents can be used	proprietary reagents —	user's option —	user's option (same capabilities when third-party reagents used) —
Walkaway capability/Walkaway duration	yes/~12.5 min. or 1 specimen or 5–6 tests	no/—	yes/22 samples plus 3 stat (reagent dependent)
Minmax. specimen volume that can be aspirated at one time Min. sample volume required for PT/PTT/Factor VIII activity	_ _	50–150 μL (total volume) 50 μL/50 μL/reagent dependent	2–275 µL 50 µL/50 µL/assay dependent
Types of disposables used	single use cartridges, weekly cleaning cartridge, QC level 1 and 2		—
Primary tube sampling supported/Pierces caps on primary tubes	yes/yes	—/no	yes/no
Accommodates most standard tube sizes/Nonstandard sizes Sample barcode-reading capability/Autodiscrimination	yes/— ves/—	no/no yes/no	yes/no yes (Interleaved 2 of 5, UPC, Codabar, Code 39, Code 128)/no
Auto tracks product volume/Measures number of tests remaining	•	no/no	yes/yes
Short sample detection	-	no	yes
Clot detection as preanalytical variable in plasma sample Auto detects adequate reagents for aspiration or analysis	_ _	no no	no yes (aspiration and analysis)
Detection or quantitation for hemolysis, turbidity, icterus, lipemia	no	detection for hemolysis, turbidity, icterus, lipemia	no
Dilutes patient samples onboard	no	no	yes
Automatic rerun capability/Auto reflex testing capability Lag time during which hypercoagulable sample not detected	no/no no	no/no no	yes/no no
User can adjust reagent volumes/Sample volumes	no/no	yes/yes	yes/yes
User can adjust No. of reagents/Sources of reagents	yes/yes	no/no	yes/yes
User can adjust incubation times/Reading times Read time extended for prolonged clotting times	no/variable	yes/yes no	yes/yes yes (selectable on menus)
Autocalibration/Calibrants stored onboard	_	no/no	yes/yes
Multipoint calibration supported/Recommended frequency	no calibration required	yes/—	yes/with lot change
Stat time to complete all analytes/Throughput per hour for:			2 minutes/120 enseimens
PT alone PT, PTT	_	_	<2 minutes/120 specimens <5 minutes/71 specimens
Fibrinogen	fibrinogen contribution to clot stiffness: ~12.5 min./—	_	<5 minutes/50 specimens
Factor VIII activity assay D-dimer			<6 minutes/— <6 minutes/—
Time delay from ordering stat to aspiration of sample	none	_	3 minutes/—
How labs get LOINC codes for results	email query, included in Quantra implementation guide, addressed during training	functionality not provided	functionality not provided
Onboard real-time QC/Onboard software capability to review QC Information that can be barcode-scanned on instrument	yes/yes operator identifier, specimen identifier, reagent lot No., lot specific QC target ranges	no/no specimen identifier, reagent lot No.	yes/yes specimen identifier
Compatible with laboratory automation systems	no onboard/	no no/	NO ophoard/
Data-management capability/LIS or EHR systems interfaced Interface supplied by instrument vendor	onboard/— yes	no/— yes (included in analyzer price)	onboard/— no
Results transferred to LIS as soon as test time complete	yes	yes	yes
Bidirectional interface capability	yes (broadcast download)	no	yes (host query)
Remote servicing provided/UPS backup power supply Instrument connections to transfer information	no/no directly to LIS; commercial middleware (RALS, Telcor, UniPOC, Cobas Infinity, DI Data Innovations)	no/no directly to LIS; directly to EHR	no/no data-management system, which in turn connects to LIS; directly to LIS
Interface standards supported Information transferred to data-management software	LIS: compliant to CLSI LIS02-A2; POC testing middleware: compliant to CLSI POCT01-A2 device unique identifier, operator ID, patient ID, result, QC	patient ID, result	LAN connection provides FTP result file transfer device unique identifier, patient ID, specimen ID, result
Avg. time for basic user training	identifier, date and time of assay start, more up to 5 days (approx. 2 days at vendor office)	1 day (at vendor office, on request)	3 days (at customer and vendor offices)
Approximate scheduled maintenance time Maintenance records kept onboard	weekly: 3 minutes; monthly: 12 minutes	per shift: <1 minute; daily: <1 minute; weekly: <1 minute; monthly: <3 minutes no	per shift: 1 minute; daily: 3 minutes; weekly: 5 minutes; monthly: 15 minutes yes
Warranty with purchase/Annual service contract cost (24/7)	yes/— (cost dependent on contract)	yes/—	yes/—
Distinguishing features (supplied by company)	ultrasound measure of whole blood hemostasis directly measures physical properties of a developing clot; wide range	advanced coagulation diagnostics by selectable dual wavelength optics (405/750 nm) for each measuring channel;	easy-to-use, standalone device with small footprint; onboard user and service software, no external PC required; optimized
Note: a dash in lieu of an answer means company did not answer question or question is not applicable	of clinical indications with simplified interpretation allows for better communication; novel technology allows for parameters not available in other systems, e.g. platelet contribution to clot	more sensitive to interferences from hemolysis, icteric, and lipemic samples; intuitive flexible user software; minimum maintenance, repair times, and costs (incl. printer)	for small to mid-sized labs
All information is supplied by the companies listed. The tabulation		, , , , , , , , , , , , , , , , , , , ,	

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Part 4 of 6	Siemens Healthineers Laura Oversmith laura.oversmith@siemens-healthineers.com Tarrytown, NY 914-631-8000 siemens-healthineers.us.com	Siemens Healthineers Laura Oversmith laura.oversmith@siemens-healthineers.com Tarrytown, NY 914-631-8000 siemens-healthineers.us.com	Siemens Healthineers Laura Oversmith laura.oversmith@siemens-healthineers.com Tarrytown, NY 914-631-8000 siemens-healthineers.us.com
Instrument name/First year sold	BCS XP Analyzer/2006	BFT II Analyzer/1999	CA-600 Series Systems/2012
List price/Model type Dimensions (H × W × D)/Weight/Instrument footprint No. of units in clinical use in U.S./Outside U.S. (countries) Composition of installs: Hospital lab/Reference lab/Other Targeted daily, monthly, annual test volume Operational type Country where analyzer designed/Manufactured	\$171,000/benchtop 37 × 49 × 25 in./330 lbs./8.5 sq. ft. >350/>1,000 (worldwide) — daily: >300 continuous random access Germany/Germany	\$8,500/benchtop 3.9 × 7.9 × 11.8 in./8.4 lbs./0.65 sq. ft. —/— (worldwide) daily: 1 batch Germany/Germany	CA-620: \$42,000; CA-660: \$55,000/benchtop 22.5 × 19.5 × 19.5 in./94.6 lbs./3.1 sq. ft. >2,000/>5,000 (worldwide) daily: <25 continuous random access Japan/Japan
Company manufactures instrument	yes	yes	no (manufactured by Sysmex)
FDA-approved clotting-based tests FDA-approved chromogenic tests	PT, APTT, fibrinogen, factors II, V, VII, VIII, IX, X, XI, XII, protein C, protein S, lupus, factor V Leiden, TT, batroxobin time antithrombin, protein C, Innovance free protein S antigen, plasminogen, Innovance heparin, factor VIII chromogenic,	PT, APTT, fibrinogen —	PT, APTT, fibrinogen, factors VII, VIII, protein C, TT, batroxobin time CA-660 only: antithrombin, protein C, Innovance heparin
FDA-approved immunologic tests	alpha-2-antiplasmin Innovance VWF Ac, Innovance D-dimer, von Willebrand factor-ristocetin cofactor assay	_	CA-660 only: Innovance D-dimer
Other FDA-approved tests User-defined tests in clinical use			
Tests in development or awaiting FDA 510(k) clearance	clot detection, optical; chromogenic; immunologic	alat detection, machanical and antical	clot detection, optical; chromogenic; immunologic
Methodologies supported Number of different measured assays onboard simultaneously Number of different assays programmed and calib. at one time No. of user-definable (open) channels/No. active simultaneously Factor assays require manual manipulation or dilutions Test throughput per hour/Assay run time Design of sample-handling system	>100 99 7,999/>100 — 380 (1 test in throughput)/1–3 min. (avg. 5 min.) continuous loading uncapped primary sample tubes and cups	clot detection, mechanical and optical 1 3/5 min. manual	5 7 7/5 — 60 (1 test in throughput)/1–3 min. (avg. 5 min.) 10-tube position sample rack
Operates on whole blood or spun plasma Reagent type Reagent barcode-reading capability	in same rack; 10-tube position sample rack spun plasma self-contained single-use vials; open reagent system (liquid, lyophilized [reconstituted manually]) yes, for all tests	spun plasma self-contained single-use vials; open reagent system (liquid, lyophilized [reconstituted manually]) no	spun plasma self-contained single-use vials; open reagent system (liquid, lyophilized [reconstituted manually]) yes, for all tests
No. of reagent containers held onboard/Reagents ready to use Reagent lot tracking/Reagent inventory/Reagents refrigerated onboard	90/yes yes/yes (15°C ± 2°C)	4/yes no/no/no	11/yes yes/yes/no (15°C ± 2°C)
Reagents, consumables loaded without interrupting testing Instrument uses proprietary or third-party reagents Maximum time same lot number of reagents can be used	yes (reagents and consumables) user's option (same capabilities when third-party reagents used) 1 year	yes (reagents and consumables) proprietary reagents, third-party reagents, user's option 1 year	no user's option (same capabilities when third-party reagents used) 1 year
Walkaway capability/Walkaway duration	yes/100 specimens or up to 400 tests	no/—	yes/10 specimens or up to 50 tests
Minmax. specimen volume that can be aspirated at one time Min. sample volume required for PT/PTT/Factor VIII activity	3 μL minimum 50 μL/50 μL/5 μL (standard)	— 50 µL/50 µL/—	5 μL minimum 50 μL/50 μL/5 μL (standard)
Types of disposables used	rotors and wash solution	cuvettes	reaction tubes, CA clean I and II
Primary tube sampling supported/Pierces caps on primary tubes	yes/no	no/no	yes/no
Accommodates most standard tube sizes/Nonstandard sizes Sample barcode-reading capability/Autodiscrimination	yes/yes yes/—	no/no no/—	yes/yes yes/—
Auto tracks product volume/Measures number of tests remaining	yes/yes	no/no	yes/no
Short sample detection Clot detection as preanalytical variable in plasma sample	yes no	no no	yes no
Auto detects adequate reagents for aspiration or analysis	yes (aspiration and analysis)	no no	no
Detection or quantitation for hemolysis, turbidity, icterus, lipemia Dilutes patient samples onboard	no yes	no no	no yes
Automatic rerun capability/Auto reflex testing capability	yes/yes	no/no	no/no
Lag time during which hypercoagulable sample not detected	yes (PT and PTT: 7 seconds)	NO	yes (PT: 7 seconds; PTT: 15 seconds)
User can adjust reagent volumes/Sample volumes User can adjust No. of reagents/Sources of reagents	yes/yes yes/yes	yes/yes yes/yes	yes/yes yes/yes
User can adjust incubation times/Reading times	yes/yes	yes/yes	yes/yes
Read time extended for prolonged clotting times Autocalibration/Calibrants stored onboard	yes yes/yes	yes no/yes	yes no/yes
Multipoint calibration supported/Recommended frequency	yes/6 months, per regulatory guidelines	yes/6 months, per regulatory guidelines	yes/6 months, per regulatory guidelines
Stat time to complete all analytes/Throughput per hour for: • PT alone • PT, PTT • Fibrinogen • Factor VIII activity assay	5 minutes/380 specimens 5 minutes/325 specimens 5 minutes/315 specimens 5 minutes/280 specimens	1 minute/1 specimen 1 minute/1 specimen	7 minutes/60 specimens 8 minutes/24 specimens 7 minutes/60 specimens 5 minutes/—
D-dimer	5 minutes/—	_	6 minutes/12 specimens
Time delay from ordering stat to aspiration of sample How labs get LOINC codes for results	1 minute website	— website	1 minute website
Onboard real-time QC/Onboard software capability to review QC Information that can be barcode-scanned on instrument Compatible with laboratory automation systems	yes/yes operator identifier, specimen identifier, reagent lot No. no	no/no — no	yes/yes specimen identifier, reagent lot No. no
Data-management capability/LIS or EHR systems interfaced Interface supplied by instrument vendor Results transferred to LIS as soon as test time complete	onboard/most major vendors contract dependent	no/no no	onboard/most major vendors contract dependent
Bidirectional interface capability	yes yes (host query)	no no	yes yes (host query)
Remote servicing provided/UPS backup power supply Instrument connections to transfer information	no/yes data-management system, which in turn connects to LIS or EHR; directly to LIS or EHR; directly to lab automation system; commercial most major companies)	no/no commercial middleware (most major companies)	no/no data-management system, which in turn connects to LIS or EHR; directly to LIS or EHR; directly to lab automation system; commercial middleware (most major companies)
Interface standards supported Information transferred to data-management software Avg. time for basic user training	ASTM 1394-91, ASTM 1381, HL7 device unique identifier, operator ID, patient ID, specimen ID, result, QC identifier, PSI checks 2 days (at customer site), some self-study virtual training required	Siemens PEPconnect online	ASTM 1394-91, ASTM 1381, HL7, CA-1000 protocol device unique identifier, patient ID, specimen ID, result, QC identifier 2 days (at customer site), some self-study virtual training required
Approximate scheduled maintenance time Maintenance records kept onboard	daily: 5 minutes; weekly: 10 minutes; monthly: 15 minutes no	daily: 1 minute no	daily: 8 minutes no
Warranty with purchase/Annual service contract cost (24/7)	yes/— (cost dependent on contract)	yes/—	yes/— (cost dependent on contract)
Distinguishing features (supplied by company)	user-definable calibration curve expiration and prewarning alerts; user-definable barcode utility enables customizable reagent protocols; user-friendly Windows 10 software	2-channel micro reagent volume clot-based technology; opto- mechanical detection accurate on lipemic, icteric samples; auto- matic INR calculation, curve storage, built-in thermal printer; effective for low-volume testing, backup to larger systems	maximizes counter space with compact footprint in low-volume labs; increases uptime and reduces service expenses; CA-620 system for routine clotting-based testing, CA-660 system for clotting, chromogenic, and immunologic testing

Note: a dash in lieu of an answer means company did not answer question or question is not applicable

Part 5 of 6	Siemens Healthineers Laura Oversmith laura.oversmith@siemens-healthineers.com Tarrytown, NY 914-631-8000 siemens-healthineers.us.com	Siemens Healthineers Laura Oversmith laura.oversmith@siemens-healthineers.com Tarrytown, NY 914-631-8000 siemens-healthineers.us.com	Werfen Catherine Quirbach cquirbach@werfen.com Bedford, MA 781-674-3221 www.werfen.com
Instrument name/First year cold			
Instrument name/First year sold	CS-2500 System/2016	CS-5100 System/2016	ACL AcuStar/2010
List price/Model type Dimensions (H × W × D)/Weight/Instrument footprint No. of units in clinical use in U.S./Outside U.S. (countries) Composition of installs: Hospital lab/Reference lab/Other Targeted daily, monthly, annual test volume Operational type Country where analyzer designed/Manufactured	\$155,000/benchtop 27 × 30.6 × 35.2 in./242.5 lbs./7.5 sq. ft. >500/>2,000 (worldwide) — daily: 25–300 continuous random access Japan/Japan	\$205,000/floor standing 50.4 × 40.6 × 45.3 in./612.9 lbs./12.8 sq. ft. >100/>1,000 (worldwide) daily: >300 continuous random access Japan/Japan	—/benchtop 21 × 34 × 24 in./170 lbs./10 sq. ft. 5/196 (available in most countries) 100%/0/0 daily: 20–150; monthly: 600–4,500; annual: 36,000–54,000 random access U.S./U.S.
Company manufactures instrument	no (manufactured by Sysmex)	no (manufactured by Sysmex)	no
FDA-approved clotting-based tests FDA-approved chromogenic tests FDA-approved immunologic tests Other FDA-approved tests	PT, APTT, fibrinogen, factors II, V, VII, VIII, IX, X, XI, XII, protein C, lupus, factor V Leiden, TT, batroxobin time antithrombin, protein C, Innovance free protein S antigen, plasminogen, Innovance heparin, factor VIII chromogenic, alpha-2-antiplasmin Innovance VWF Activity, VWF Antigen, Innovance D-dimer	PT, APTT, fibrinogen, factors II, V, VII, VIII, IX, X, XI, XII, protein C, lupus, factor V Leiden, TT, batroxobin time antithrombin, protein C, Innovance free protein S antigen, plasminogen, Innovance heparin, factor VIII chromogenic, alpha-2-antiplasmin Innovance VWF Activity, VWF Antigen, Innovance D-dimer	— — — — HIT IgG, domain 1, anticardiolipin IgG, anticardiolipin IgM, B2GPI IgG, B2GPI IgM
User-defined tests in clinical use Tests in development or awaiting FDA 510(k) clearance	platelet aggregation RUO Innovance Anti-Xa assay	platelet aggregation RUO Innovance Anti-Xa assay	ADAMTS13, von Willebrand factor ristocetin cofactor, von
Methodologies supported	clot detection, optical; chromogenic; immunologic	clot detection, optical; chromogenic; immunologic	Willebrand antigen immunologic (chemiluminescent)
Number of different measured assays onboard simultaneously Number of different assays programmed and calib. at one time	60 60 00 000/00	60 60	20 20 20
No. of user-definable (open) channels/No. active simultaneously Factor assays require manual manipulation or dilutions	80,000/60 —	80,000/60 	0/0
Test throughput per hour/Assay run time Design of sample-handling system Operates on whole blood or spun plasma	180 (2 tests in throughput)/1–10 min. (avg. 5 min.) continuous loading capped and uncapped primary sample tubes and cups in same rack; 10-tube position sample rack \times 5 spun plasma	400 (2 tests in throughput)/1–10 min. (avg. 5 min.) continuous loading capped and uncapped primary sample tubes and cups in same rack; 10-tube position sample rack × 10 spun plasma	60 (1 test in throughput)/30 min. samples loaded into carousel rack spun plasma
Reagent type	self-contained single-use vials; open reagent system (liquid, lyophilized [reconstituted manually])	self-contained single-use vials; open reagent system (liquid, lyophilized [reconstituted manually])	self-contained multiuse cartridges (liquid)
Reagent barcode-reading capability No. of reagent containers held onboard/Reagents ready to use	yes, for all tests 40/yes	yes, for all tests 40/yes	yes, for all tests 20/yes
Reagent lot tracking/Reagent inventory/Reagents refrigerated onboard	yes/yes (10°C ± 2°C)	yes/yes/yes (10°C ± 2°C)	yes/yes/yes
Reagents, consumables loaded without interrupting testing Instrument uses proprietary or third-party reagents Maximum time same lot number of reagents can be used	yes (reagents and consumables) user's option (same capabilities when third-party reagents used) 1 year	yes (reagents and consumables) user's option (same capabilities when third-party reagents used) 1 year	yes (reagents and consumables) proprietary reagents 6 months
Walkaway capability/Walkaway duration	yes/50 specimens or up to 500 tests	yes/100 specimens or up to 1,000 tests	yes/30 specimens or 280 tests
Minmax. specimen volume that can be aspirated at one time Min. sample volume required for PT/PTT/Factor VIII activity	5 μL minimum 50 μL/50 μL/5 μL (standard)	5 μL minimum 50 μL/50 μL/5 μL (standard)	2–250 μL 50 μL/50 μL/50 μL
Types of disposables used	reaction tubes, CA clean I and II	reaction tubes, CA clean I and II	cuvettes, trigger, solutions
Primary tube sampling supported/Pierces caps on primary tubes	yes/yes	yes/yes	yes/no
Accommodates most standard tube sizes/Nonstandard sizes	yes/yes	yes/yes	yes/no
Sample barcode-reading capability/Autodiscrimination Auto tracks product volume/Measures number of tests remaining	yes/— yes/yes	yes/— yes/yes	yes/yes yes/yes
Short sample detection	yes	yes	yes
Clot detection as preanalytical variable in plasma sample	yes	yes	no
Auto detects adequate reagents for aspiration or analysis Detection or quantitation for hemolysis, turbidity, icterus, lipemia	yes (aspiration and analysis) detection and quantitation for hemolysis, turbidity, icterus, lipemia	yes (aspiration and analysis) detection and quantitation for hemolysis, turbidity, icterus, lipemia	yes (aspiration and analysis)
Dilutes patient samples onboard	yes	yes	yes
Automatic rerun capability/Auto reflex testing capability	yes/yes	yes/yes	yes/yes
Lag time during which hypercoagulable sample not detected User can adjust reagent volumes/Sample volumes	yes (PT: 7 seconds; PTT: 15 seconds) yes/yes	yes (PT: 7 seconds; PTT: 15 seconds) yes/yes	no/no
User can adjust No. of reagents/Sources of reagents	yes/yes	yes/yes	no/no
User can adjust incubation times/Reading times	yes/yes	yes/yes	no/no
Read time extended for prolonged clotting times Autocalibration/Calibrants stored onboard	yes (selectable on menus) no/yes	yes (selectable on menus) no/yes	no no/no
Multipoint calibration supported/Recommended frequency	yes/6 months, per regulatory guidelines	yes/6 months, per regulatory guidelines	yes/6 months
Stat time to complete all analytes/Throughput per hour for: • PT alone • PT, PTT • Fibrinogen	5 minutes/180 specimens 5 minutes/90 specimens 5 minutes/180 specimens	5 minutes/400 specimens 5 minutes/200 specimens 5 minutes/201 specimens	_ _ _
Factor VIII activity assay D-dimer	5 minutes/— 5 minutes/90 specimens	5 minutes/— 5 minutes/202 specimens	
Time delay from ordering stat to aspiration of sample	1 minute	1 minute	<1 minute
How labs get LOINC codes for results	website	website	functionality not provided
Onboard real-time QC/Onboard software capability to review QC Information that can be barcode-scanned on instrument	yes/yes operator identifier, specimen identifier, reagent lot No.	yes/yes operator identifier, specimen identifier, reagent lot No.	yes/yes specimen identifier, reagent lot No.
Compatible with laboratory automation systems	no	yes (Siemens Aptio Automation)	no
Data-management capability/LIS or EHR systems interfaced	onboard/most major vendors contract dependent	onboard/most major vendors contract dependent	onboard/Meditech
Interface supplied by instrument vendor Results transferred to LIS as soon as test time complete	yes	yes	contract dependent yes
Bidirectional interface capability	yes (host query)	yes (host query)	yes (broadcast download and host query)
Remote servicing provided/UPS backup power supply Instrument connections to transfer information	yes/yes data-management system, which in turn connects to LIS or EHR; directly to LIS or EHR; directly to lab automation system; commercial middleware (most major companies)	yes/yes data-management system, which in turn connects to LIS or EHR; directly to LIS or EHR; directly to lab automation system; commercial middleware (most major companies)	no/yes data-management system, which in turn connects to LIS; directly to LIS; commercial middleware (Werfen, Beckman Coulter)
Interface standards supported Information transferred to data-management software	ASTM 1394-91, ASTM 1381 device unique identifier, operator ID, patient ID, specimen ID, result, QC identifier, PSI checks	ASTM 1394-91, ASTM 1381 device unique identifier, operator ID, patient ID, specimen ID, result, QC identifier, PSI checks	ASTM 1394-91 specimen ID
Avg. time for basic user training Approximate scheduled maintenance time Maintenance records kept onboard	2 days (at customer site), some self-study virtual training required daily: 5 minutes; weekly: 1 minute; monthly: 1 minute yes	2 days (at customer site), some self-study virtual training required daily: 5 minutes; weekly: 1 minute; monthly: 1 minute yes	4 days (at customer site) daily: 5 minutes; weekly: 5 minutes yes
Warranty with purchase/Annual service contract cost (24/7)	yes/— (cost dependent on contract)	yes/— (cost dependent on contract)	yes/— (cost dependent on contract)
Distinguishing features (supplied by company)	confidence: simultaneous multiwavelength detection and PSI technology designed to ensure high-quality first-run results; operational efficiency: user-friendly software on Windows 10; tilted reagent vials for efficiency; consistency: for multisite	confidence: simultaneous multiwavelength detection and PSI technology designed to ensure high-quality first-run results; operational efficiency: user-friendly software on Windows 10; tilted reagent vials for efficiency; consistency: for multisite	on-demand HIT IgG testing with results available in 30 minutes; uses sensitive chemiluminescent technology, improving sensitivity; reagents are ready to use with onboard stability up to 12 weeks
Note: a dash in lieu of an answer means company did not answer question or question is not applicable	patient monitoring, with sample result traceability for in-depth audit capabilities	patient monitoring, with sample result traceability for in-depth audit capabilities	

Part 6 of 6	Werfen Catherine Quirbach cquirbach@werfen.com Bedford, MA 781-674-3221 www.werfen.com	Werfen Catherine Quirbach cquirbach@werfen.com Bedford, MA 781-674-3221 www.werfen.com	Werfen Catherine Quirbach cquirbach@werfen.com Bedford, MA 781-674-3221 www.werfen.com
Instrument name/First year sold	ACL TOP 350/2016	ACL TOP 550 CTS/2016	ACL TOP 750 Series/2016
List price/Model type Dimensions (H × W × D)/Weight/Instrument footprint No. of units in clinical use in U.S./Outside U.S. (countries) Composition of installs: Hospital lab/Reference lab/Other Targeted daily, monthly, annual test volume Operational type Country where analyzer designed/Manufactured Company manufactures instrument	—/benchtop 29 × 32 × 33 in./200 lbs./8 sq. ft. 1,309/3,700 (available in most countries) 95%/5%/— daily: 20–150; monthly: 600–4,500; annual: 36,000–54,000 random access U.S./U.S. yes	—/benchtop 29 × 43 × 35 in./312 lbs./14 sq. ft. 737/3,400 (available in most countries) 85%/12%/3% (pharmaceutical or medical device labs) daily: 100–200; monthly: 3,000–6,000; annual: 36,000–72,000 random access U.S./U.S. yes	—/floor standing 29 × 60 × 35 in./356 lbs./21 sq. ft. 737/3,200 (available in most countries) 85%/12%/3% (pharmaceutical or medical device labs) daily: 200–400; monthly: 6,000–12,000; annual: 72,000–144,000 random access U.S./U.S. yes
FDA-approved clotting-based tests	PT, APTT, fibrinogen, thrombin time, factor assays, lupus	PT, APTT, fibrinogen, thrombin time, factor assays, lupus	PT, APTT, fibrinogen, thrombin time, factor assays, lupus
FDA-approved chromogenic tests FDA-approved immunologic tests Other FDA-approved tests	anticoagulant (dRWT and silica clotting time), protein S, protein C anti-Xa, apixaban, protein C, antithrombin, plasminogen, plasmin inhibitor high specificity D-dimer, standard D-dimer, heparin induced thrombocytopenia, von Willebrand factor antigen, von Willebrand factor activity, free protein S, factor XIII antigen, homocysteine	anticoagulant (dRWT and silica clotting time), protein S, protein C anti-Xa, apixaban, protein C, antithrombin, plasminogen, plasmin inhibitor high specificity D-dimer, standard D-dimer, heparin induced thrombocytopenia, von Willebrand factor antigen, von Willebrand factor activity, free protein S, factor XIII antigen, homocysteine	anticoagulant (dRWT and silica clotting time), protein S, protein C anti-Xa, apixaban, protein C, antithrombin, plasminogen, plasmin inhibitor high specificity D-dimer, standard D-dimer, heparin induced thrombocytopenia, von Willebrand factor antigen, von Willebrand factor activity, free protein S, factor XIII antigen, homocysteine
User-defined tests in clinical use	DOAC assays, chromogenic factor VIII	DOAC assays, chromogenic factor VIII	DOAC assays, chromogenic factor VIII
Tests in development or awaiting FDA 510(k) clearance	von Willebrand factor ristocetin cofactor, dabigatran, rivaroxaban, apixaban	von Willebrand factor ristocetin cofactor, dabigatran, rivaroxaban, apixaban	von Willebrand factor ristocetin cofactor, dabigatran, rivaroxaban, apixaban
Methodologies supported	clot detection, optical; chromogenic; immunologic	clot detection, optical; chromogenic; immunologic	clot detection, optical; chromogenic; immunologic
Number of different measured assays onboard simultaneously Number of different assays programmed and calib. at one time No. of user-definable (open) channels/No. active simultaneously Factor assays require manual manipulation or dilutions	(immunoturbidimetric) 30 500 250/250	(immunoturbidimetric) 30 500 250/250	(immunoturbidimetric) 30 500 250/250
Test throughput per hour/Assay run time Design of sample-handling system Operates on whole blood or spun plasma	110 (1 test in throughput)/3–6 min. (avg. 4 min.) samples loaded into rack; system uses integrated sample barcode reader to eliminate any post-draw sample misidentification spun plasma	240 (1 test in throughput)/3–6 min. (avg. 4 min.) samples loaded into rack; system uses integrated sample barcode reader to eliminate any post-draw sample misidentification spun plasma	360 (1 test in throughput)/3–6 min. (avg. 4 min.) samples loaded into rack; system uses integrated sample barcode reader to eliminate any post-draw sample misidentification spun plasma
Reagent type Reagent barcode-reading capability No. of reagent containers held onboard/Reagents ready to use	self-contained multiuse vials; open reagent system (liquid, lyophilized [reconstituted manually]) yes, for all tests 24/variable (reagent specific)	self-contained multiuse vials; open reagent system (liquid, lyophilized [reconstituted manually]) yes, for all tests 40/variable (reagent specific)	self-contained multiuse vials; open reagent system (liquid, lyophilized [reconstituted manually]) yes, for all tests 60/variable (reagent specific)
Reagent lot tracking/Reagent inventory/Reagents refrigerated onboard Reagents, consumables loaded without interrupting testing Instrument uses proprietary or third-party reagents	yes/yes/yes yes (reagents and consumables) user's option (same capabilities when third-party reagents used)	yes/yes/yes yes (reagents and consumables) user's option (same capabilities when third-party reagents used)	yes/yes/yes yes (reagents and consumables) user's option (same capabilities when third-party reagents used)
Maximum time same lot number of reagents can be used	18 months	18 months	18 months
Walkaway capability/Walkaway duration	yes/40 specimens or 800 tests	yes/80 specimens or 800 tests	yes/120 specimens or 800 tests
Min.—max. specimen volume that can be aspirated at one time	2–250 μL	2–250 μL	2–250 μL
Min. sample volume required for PT/PTT/Factor VIII activity Types of disposables used	50 μL/50 μL/50 μL cuvettes, clean A/B, rinse	50 μL/50 μL/50 μL cuvettes, clean A/B, rinse	50 μL/50 μL/50 μL cuvettes, clean A/B, rinse
Primary tube sampling supported/Pierces caps on primary tubes	yes/yes	yes/yes	yes/yes
Accommodates most standard tube sizes/Nonstandard sizes Sample barcode-reading capability/Autodiscrimination	yes/yes yes (Interleaved 2 of 5, Code 39, Code 128)/yes	yes/yes yes (Interleaved 2 of 5, Code 39, Code 128)/yes	yes/yes yes (Interleaved 2 of 5, Code 39, Code 128)/yes
Auto tracks product volume/Measures number of tests remaining	yes/yes	yes/yes	yes/yes
Short sample detection Clot detection as preanalytical variable in plasma sample	yes yes	yes yes	yes yes
Auto detects adequate reagents for aspiration or analysis Detection or quantitation for hemolysis, turbidity, icterus, lipemia Dilutes patient samples onboard	yes (aspiration and analysis) detection and quantitation for hemolysis, turbidity, icterus, lipemia yes	yes (aspiration and analysis) detection and quantitation for hemolysis, turbidity, icterus, lipemia yes	yes (aspiration and analysis) detection and quantitation for hemolysis, turbidity, icterus, lipemia yes
Automatic rerun capability/Auto reflex testing capability Lag time during which hypercoagulable sample not detected	yes/yes no	yes/yes no	yes/yes no
User can adjust reagent volumes/Sample volumes	yes/yes	yes/yes	yes/yes
User can adjust No. of reagents/Sources of reagents	yes/yes	yes/yes	yes/yes
User can adjust incubation times/Reading times Read time extended for prolonged clotting times	yes/yes yes (selectable on menus)	yes/yes yes (selectable on menus)	yes/yes yes (selectable on menus)
Autocalibration/Calibrants stored onboard	no/no	no/no	no/no
Multipoint calibration supported/Recommended frequency	yes/6 months	yes/6 months	yes/6 months
Stat time to complete all analytes/Throughput per hour for: • PT alone • PT, PTT • Fibrinogen	<3 minutes/110 specimens <6 minutes/55 specimens <3 minutes/60 specimens	<3 minutes/240 specimens <6 minutes/90 specimens <3 minutes/78 specimens	<3 minutes/360 specimens <6 minutes/165 specimens <3 minutes/108 specimens
Factor VIII activity assay D-dimer	8 minutes/38 specimens 5 minutes/55 specimens	8 minutes/77 specimens 5 minutes/75 specimens	8 minutes/100 specimens 5 minutes/100 specimens
Time delay from ordering stat to aspiration of sample How labs get LOINC codes for results	none functionality not provided	none functionality not provided	none functionality not provided
Onboard real-time QC/Onboard software capability to review QC Information that can be barcode-scanned on instrument Compatible with laboratory automation systems	yes/yes specimen identifier, reagent lot No.	yes/yes specimen identifier, reagent lot No.	yes/yes specimen identifier, reagent lot No. yes (HemoCell, Beckman, Siemens, Abbott, Thermo Fisher, others)
Data-management capability/LIS or EHR systems interfaced Interface supplied by instrument vendor	no onboard/Cerner, CompuLab, CPSI, HBOC, HMS, McKesson, Meditech, Novius, SMS, SCC, Clinisys, Vista contract dependent	no onboard/Cerner, CompuLab, CPSI, HBOC, HMS, McKesson, Meditech, Novius, SMS, SCC, Clinisys, Vista contract dependent	onboard/Cerner, CompuLab, CPSI, HBOC, HMS, McKesson, Meditech, Novius, SMS, SCC, Clinisys, Vista contract dependent
Results transferred to LIS as soon as test time complete Bidirectional interface capability	yes yes (broadcast download and host query)	yes yes (broadcast download and host query)	yes yes (broadcast download and host query)
Remote servicing provided/UPS backup power supply Instrument connections to transfer information	yes/yes data-management system, which in turn connects to LIS; directly to LIS; commercial middleware (Beckman Coulter)	yes/yes data-management system, which in turn connects to LIS; directly to LIS; commercial middleware (Beckman Coulter)	yes/yes data-management system, which in turn connects to LIS; directly to LIS; directly to lab automation system; commercial middleware (Beckman Coulter)
Interface standards supported Information transferred to data-management software Avg. time for basic user training Approximate scheduled maintenance time Maintenance records kept onboard	ASTM 1394-91 specimen ID 9 days (5 days at customer site, 4 days at vendor office) daily: <5 minutes; weekly: <10 minutes yes	ASTM 1394-91 specimen ID 14 days (10 days at customer site, 4 days at vendor office) daily: <5 minutes; weekly: <10 minutes yes	ASTM 1394-91 specimen ID 14 days (10 days at customer site, 4 days at vendor office) daily: <5 minutes; weekly: <10 minutes yes
Warranty with purchase/Annual service contract cost (24/7) Distinguishing features (supplied by company)	yes/— (cost dependent on contract) FDA 510(k) approved; improves patient care, lab efficiencies,	yes/— (cost dependent on contract) FDA 510(k) approved; improves patient care, lab efficiencies,	yes/— (cost dependent on contract) FDA 510(k) approved; improves patient care, lab efficiencies,
Note: a dash in lieu of an answer means company did not answer question or question is not applicable	and costs; supports lab's policy on sample acceptance with quantitative HIL, sample fill detection, aspiration errors (clots); best-in-class reagents; liquid format for PT/APTT (10-day onboard stability), high specificity D-dimer, on-demand HIT testing	and costs; supports lab's policy on sample acceptance with quantitative HIL, sample fill detection, aspiration errors (clots); liquid format for PT/APTT (10-day onboard stability), high specificity D-dimer, on-demand HIT testing	and costs; supports lab's policy on sample acceptance with quantitative HIL, sample fill detection, aspiration errors (clots); liquid format for PT/APTT (10-day onboard stability), high specificity D-dimer, on-demand HIT testing