

Definition of efficient multiplex diagnostics

Microblot-Array is an immunoblot array in microtiter plate format designed for efficient multiplex diagnostics. The technology eliminates the bottleneck of traditional BLOT processing and capacity and opens up the way to high throughput testing and automation.

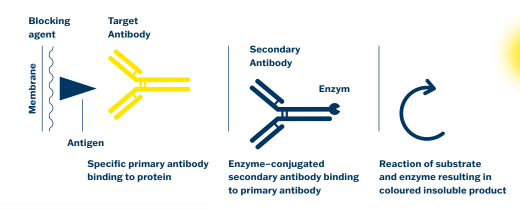
The comprehensive evaluation of Microblot Array testing is ensured by using the Microblot-Array Software in combination with the Microblot-Array Reader, enabling complex image analysis including results evaluation and connectivity to LIS.

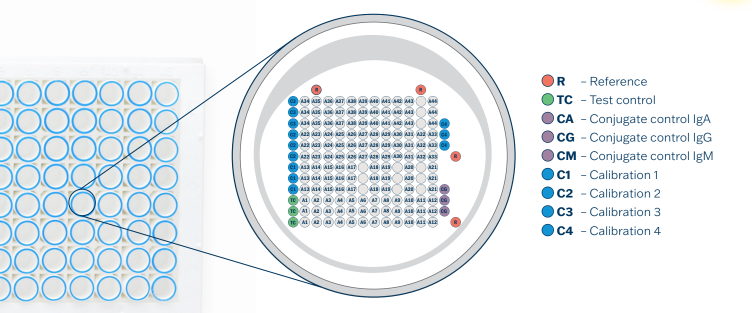
Main clinical areas covered

- Infectious serology
- Autoimmunity

Microblot-Array principle

Specific recombinant proteins/antigens spotted onto a nitrocellulose membrane





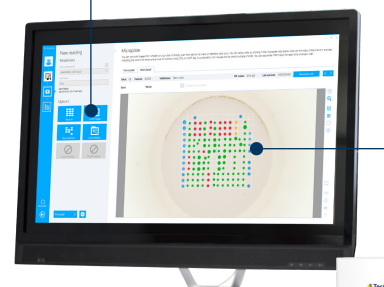
Complex solution

Microblot-Array Software

- Automated test identification
- Intuitive and user-friendly guiding throughout the results evaluation
- Complex image analysis
- Optional manual control of spot localization
- Detailed results comparison within single wells and spots
- Evaluation of the validity test through control spots
- Export of results in various formats
- LIS connectivity

Calibration data

- Innovative processing and evaluation with LOT identification
- Calibration data ensure significant benefits:
 - Interchangeability of conjugate and substrate between the same Ig classes
 - Unification of evaluation criteria for all MBA kits
 - The more effective automatic processing



Microblot-Array

- Antigens spotted in triplicate minimizing statistical variation
- Controls in each well
- 4 calibration spots to create a calibration curve
- Evaluation based on combination of positive antigen spots: qualitative, quantitative (U/ml)
- or semiquantitative (IP)

Microblot-Array Reader

- Fast high-quality scanning and evaluation: 5 min. per full plate
- Scanning of selected wells
- Automated spot localization and image analysis
- Optimized for a 96-well microtiter plates format

Benefits

Efficiency

- Analysis of up to 96 patient samples per plate
- Low sample consumption 10 μl
- Parallel testing of multiple markers simultaneously
- Time and cost saving diagnostics

Flexibility

- One parameter × various parameters
- One well × high number of samples
- Manual processing × automated processing

Automation

- Possibility of automated processing using an ELISA instrument
- Intuitive software for test evaluation
- Evaluation of individual antigens and their association with pathogen species or disease type

User comfort

- Ready-to-use components
- Identical assay procedure (30-30-15 min.)
- Remote troubleshooting
- Reagents interchangeability due to batch identification (calibration data)



Automatic processing by ELISA analyzer minimizes hands-on time, eliminates errors rate due to the QR code identification system, and improves the throughput of samples.



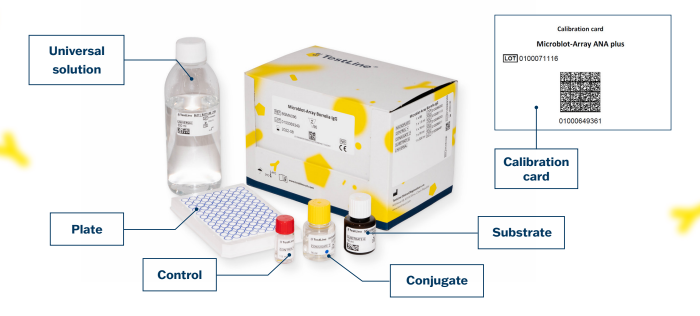
Protocol Summary

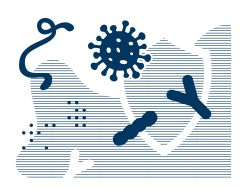
Step No. Test steps

- **1.** Pipette Universal Solution 150 μl
- **2.** Wells soaking at room temperature for 10 min.
- **3.** Aspirate off

Dilute samples

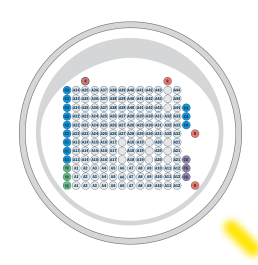
- 4. serum/plasma 1:51 (10 μl + 500 μl) cerebrospinal fluid 1:3 (50 μl + 100 μl) synovial fluid 1:17.5 (10 μl + 165 μl)
- **5.** Pipette control and diluted samples 100 μl
- **6.** Incubate at room temperature for 30 min.
- **7.** Quick wash using the Universal Solution
- 8. Aspirate and wash 3 × 5 min. with 150 μl of Universal Solution
- 9. Pipette Conjugate 100 μl
- **10.** Incubate at room temperature for 30 min.
- **11.** Quick wash using the Universal Solution
- **12.** Aspirate and wash 3 × 5 min. with 150 μl of Universal Solution
- **13.** Pipette Substrate Solution (BCIP/NBT) 100 μl
- **14.** Incubate at room temperature for 15 min.
- **15.** Quick wash using the distilled water
- **16.** Aspirate and wash 2 × 5 min. with 200 μl of distilled water
- **17.** Dry and evaluate strips





Microblot-Array for the diagnostics of systemic autoimmune diseases

The main benefit of Microblot-Array ANA kits is the high number of antigens which can be simultaneously detected in one sample. The kits are primarily intended for confirmation of ELISA or other screening method. However, they also enable identification of specific antibody and thus differentiation of systemic autoimmune diseases, such as myositis, scleroderma, systemic lupus and others. The kits are optimized and validated for detection of specific IgG in human serum or plasma.



Test Characteristics

Parameters of the Microblot-Array ANA kit

| | Diagnostic Sensitivity | Diagnostic Specificity |
|-----|------------------------|-------------------------------|
| ANA | 95.2% (n = 398) | 95.3% (n = 148) |

Comparative Study - Correlation of Results

Myopathy

| <u>n = 80</u> | Microblot-Array | BLOT |
|------------------|-----------------|-------------|
| positive | 70 | 69 |
| negative | 0 | 0 |
| total conformity | 98.6 % | |

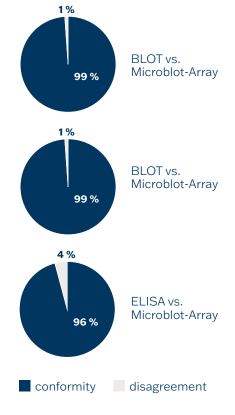
Systemic sclerosis

total conformity

| <u>n = 124</u> | Microblot-Array | BLOT | |
|----------------|-----------------|------|--|
| positive | 107 | 106 | |
| negative | 0 | Ο | |

99.1 %

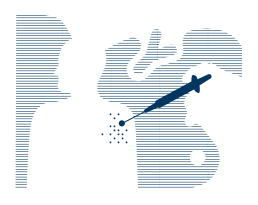
| <u>n = 204</u> | Microblot-Array | ELISA |
|------------------|-----------------|-------|
| positive | 194 | 186 |
| negative | 7 | 0 |
| total conformity | 95.5 | % |



| Spot No. | Antigen | Description |
|----------|---------|-------------|

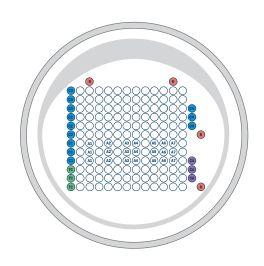
| | | | | | ma | SLE and other connective tissue diseases |
|------------|---------------|--|-----|----------|-------------|---|
| | | | | sitis | Scleroderma | SLE and other connective tissue disease |
| | | | ANA | Myositis | Scler | SLE |
| A1 | Jo-1 | Hystidyl tRNA synthetase | | | | |
| A2 | PL-7 | Threonyl tRNA synthetase | | | | |
| A3 | PL-12 | Alanyl tRNA synthetase | | | | |
| A4 | EJ | Glycyl tRNA Synthetase | • | • | | |
| A5 | OJ | IsoleucyI tRNA synthetase | • | • | | |
| A6 | KS | Asparaginyl tRNA synthetase | • | • | | |
| A7 | YARS | Tyrosyl tRNA synthetase (Ha) | • | • | | |
| A8 | ZoA | Phenylalanyl tRNA synthetase | • | • | | |
| A9 | ZoB | Phenylalanyl tRNA synthetase | • | • | | |
| A10 | HMGCR* | 3-hydroxy-3methylglutaryl-coenzyme A reductase | • | • | | |
| A11 | SAE-1 | Small ubiquitin-like modifier activating enzyme | • | • | | |
| A12 | SAE-2 | Small ubiquitin-like modifier activating enzyme | • | • | | |
| A13 | SRP54 | Signal recognition particle | • | • | | |
| A14 | Mi-2 | Helicase protein-nuclear transcription | • | • | | |
| A15 | TIF1γ | Transcription Intermediary Factor 1 | • | • | | |
| A16 | MDA5 | Melanoma differentiation associated protein 5 | • | • | | |
| | | (CADM-140) | - | - | | |
| A17 | NXP2 | Nuclear matrix protein 2 (p140, MJ) | • | • | | |
| A18 | PMScl 100 | Human exosome complex | • | • | • | |
| A19 | PMScl 75 | Human exosome complex | • | • | • | |
| A20 | ScI70 | DNA-topoisomerase I | • | | • | |
| A21 A22 | CENP A CENP B | Centromere A Centromere B | • | | • | |
| A23 | POLR3A | | • | | • | |
| A23 | NOR90 | RNA polymerase III Nucleolar transcription factor 1 (Ubtf1) | | | • | |
| A25 | Th/To | Ribonuclease P protein subunit 25 (Rpp25) | | | | • |
| A26 | PDGFR-β | Platelet-derived growth factor receptor beta | | | | |
| A27 | Fibrillarin | U3 RNP - fibrillarin | | | | |
| A28 | Ro52 | TRIM21 | | | | |
| A29 | Ro60 | Sjögren's-syndrome-related antigen A (SS-A) | • | | | • |
| A30 | La | Sjögren's-syndrome-related antigen B (SS-B) | • | | | • |
| A31 | RNPA | U1 small nuclear ribonucleoprotein A | • | | • | • |
| A32 | RNP 68/70 | U1 small nuclear ribonucleoprotein 68/70 kDa | • | | • | • |
| A33 | RNPC | U1 small nuclear ribonucleoprotein C | • | | • | • |
| A34 | SmB | Smith antigen B | • | | | • |
| A35 | SmD | Smith antigen D | • | | | • |
| A36 | PCNA | Proliferating cell nuclear antigen | • | | | • |
| A37 | P0 | Ribosomal protein P0 | • | | | • |
| A38 | Ku | Ku (p70/p80) | • | • | • | • |
| A39 | Nucleolin | Nucleolin | • | | | • |
| A40 | Histons | Histone | • | | | • |
| A41 | Nucleosome | Nucleosome | • | | | • |
| A42 | dsDNA | Double-stranded DNA | • | | | • |
| A43 | M2 | Mitochondrial M2 (AMA-M2) | • | | • | |
| A44 | DFS70 | Dense fine speckled 70 antigen | • | | | |

^{*}Check availability in your country.
• - supplementary antigens, SLE - Systemic lupus erythematosus



Microblot-array for the diagnostics of Bordetella pertussis and Bordetella parapertussis

Microblot-Array Bordetella kits provide the detailed determination of the presence of specific IgA, IgG, and IgM antibodies to recombinant Bordetella pertussis and Bordetella parapertussis antigens in human serum or plasma. It can be used for differentiation of postinfection and postvaccination antibodies as well as for differentiation disease stage. It confirms positive or borderline ELISA or agglutination test.



Test Characteristics

Parameters of Microblot-Array Bordetella kits

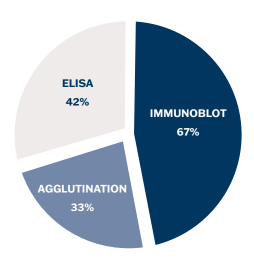
| Pathogen | Diagnostic Sensitivity | Diagnostic Specificity |
|--|------------------------|-------------------------------|
| Microblot-Array Bordetella pertussis IgA | 95.4% | 100.0% |
| Microblot-Array Bordetella parapertussis IgA | 96.9% | 100.0% |
| Microblot-Array Bordetella pertussis IgG | 97.6% | 100.0% |
| Microblot-Array Bordetella parapertussis IgG | 97.1% | 100.0% |
| Microblot-Array Bordetella pertussis IgM | 95.4% | 100.0% |
| Microblot-Array Bordetella parapertussis IgM | 95.8% | 100.0% |



| Spot No. | <u>Antigen</u> | Description | Pathogen |
|----------|----------------|---|-------------------------|
| A1 | PT | Pertussis toxin (45 kDa) – basic virulence factor, specific only for <i>B. pertussis</i> , the most important pertussis antigen | |
| A2 | FHA | B. pertussis filamentous hemagglutinin – adhesive protein, important immunogen; selected part of the sequence with high specificity | Davidatalla |
| A3 | ACT | Adenylate cyclase toxin (CyaA) – significant virulence factor of <i>B. pertussis</i> with anti-phagocytic activity | Bordetella pertussis |
| A4 | TCF | Tracheal colonization factor – protein produced only by B. pertussis; adhesin; enabling the microorganism to adhere to mucosal surfaces of respiratory tract and colonize ciliated epithelial cells and phagocytes | |
| A5 | Pertactin | 75 kDa; outer membrane protein of virulent <i>B. parapertussis</i> strains | Bordetella |
| A6 | FimN | Fimbriae N – adhesin, non-produced by <i>B. pertussis</i> | parapertussis |
| A7 | EntA | Entericidin A – membrane lipoprotein | |

Clinical Data

Laboratory detection of acute infection in a group of patients with clinical diagnosis of pertussis (n= 25 paired samples)

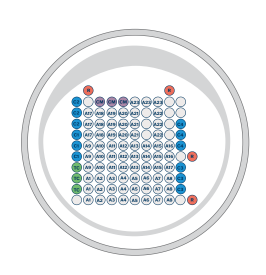




Microblot-Array for the diagnostics of Borrelia species and Anaplasma phagocytophilum

The kits are optimized for the detection of specific IgG and IgM antibodies to recombinant antigens of Borrelia species and Anaplasma phagocytophilum (HGA) in human serum, plasma, cerebrospinal or synovial fluid.

Serological diagnostics of borreliosis is difficult due to the large genetic diversity of the species Borrelia burgdorferi s.l., possible cross reactivity with unrelated antigens of other microorganisms (p44, OmpA, TpN17 and VCA-p18), and borrelia richness to heat shock proteins. Diagnostics is also complicated due to various individual serological reactivity. The production of antibodies can be extremely slow in the early phase of the disease. On the other hand, the IgG and IgM antibodies can persist for more than ten years. The Microblot-Array Borrelia kits help to refine the diagnostics thanks to the high number of antigens present in one single test.



Test characteristics

Parameters of Microblot-Array Borrelia IgG (tested on sera)

| | Diagnostic | Diagnostic |
|---------------|----------------|-----------------|
| | Sensitivity | Specificity |
| Borrelia IgG | 97.3% (n = 74) | 98.0% (n = 100) |
| Anaplasma IgG | 92.0% (n = 25) | 100.0% (n = 30) |
| Treponema | 98.3% (n = 59) | 100.0% (n = 30) |

Parameters of Microblot-Array Borrelia IgM (tested on sera)

| | Diagnostic | Diagnostic | |
|---------------|-----------------|-----------------|--|
| | Sensitivity | Specificity | |
| Borrelia IgM | 94.6% (n = 56) | 95.8% (n = 95) | |
| Anaplasma IgM | 95.0% (n = 20) | 100.0% (n = 38) | |
| EBV | 100.0% (n = 39) | 98.0% (n = 51) | |

Comparative Study

Correlation of results IgG

| <u>n = 77</u> | Microblot-Array | ELISA |
|------------------|-----------------|-------|
| positive | 38 | 41 |
| negative | 33 | 36 |
| total conformity | 92.2 | % |

Correlation of results IgM

| <u>n = 68</u> | Microblot-Array | ELISA |
|------------------|-----------------|-------|
| positive | 19 | 21 |
| negative | 40 | 44 |
| total conformity | 90.79 | V. |

8 % 92%

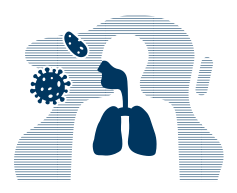
ELISA vs. Microblot-Array IgG ELISA vs. Microblot-Array IgM

conformity disagreement

| Spot No. | Antigen | Description | Kit |
|--------------------------|---|---|---------------------------------|
| A1 A2 A3 | VIsE Ba VIsE Bg VIsE Bs | Expressed part of variable major protein-like sequence, significant for IgG antibody response, species-specific antigen | |
| A4 | p83 | Main extracellular protein (product of p100 degradation) | |
| A5 | p58 | OppA-2 (Oligopeptide permease 2) – membrane transporter, is considered a marker of disseminated stage of Lyme disease | |
| A6 A7 | p41 Ba p41 Bs | Internal flagellin, highly specific antigen of early antibody response | |
| A8 | p39 | BmpA (glycosaminopeptide receptor) – marker of late IgG antibody response | |
| A9 | OspB | Outer surface protein B, marker of late stage of infection, considered a marker of Lyme arthritis | |
| A10 A11 A12 | OspA Ba OspA Bg OspA Bs | Outer surface protein A, highly specific marker of <i>Borrelia</i> infection in IgG class | Microblot-Array Borrelia IgG |
| A13 A14 A15 A16 | OspC Ba OspC Bg OspC Bs OspC Bsp | Outer surface protein C – main antigen of early antibody response, immunodominant marker of IgM antibody response | Microblot-Array Borrelia IgM |
| A17 | OspE | Outer surface protein E | |
| A18 | NapA | Neutrophil activating protein A – strong immunogen, main marker of Lyme arthritis pathogenesis | |
| A19 | p17 | DbpA (decorin-binding protein A) – outer membrane protein | |
| A20 | p44 | Anaplasma phagocytophilum – main marker of HGA antibody response | |
| A21 | OmpA | Outer membrane protein A of <i>Anaplasma phagocytophilum</i> ; peptidoglycan-associated lipoprotein, significant virulence marker | |
| A22 | Asp62 | Surface protein – membrane transporter | |
| A23 | TpN17 | Highly specific membrane protein of Treponema pallidum | Microblot-Array Borrelia IgG |
| 7.25 | VCA-p18 | Viral Capsid Antigen p18 – important marker of EBV infection | Microblot-Array Borrelia IgM |

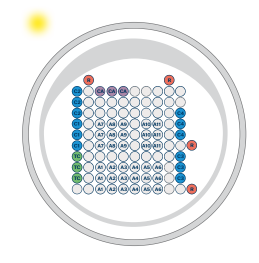
(Ba – B. afzelii, Bg – B. garinii, Bs – B. burgdorferi sensu stricto, Bsp – B. spielmanii)





Microblot-Array for the diagnostics of *Chlamydia* species

Microblot-Array Chlamydia are kits designed for the confirmation of positive or cut-off results of samples which were previously screened by ELISA or other serological methods. They serve for the detection of specific IgA and IgG antibodies to recombinant antigens of *Chlamydia* species in human serum or plasma. Thanks to the complex antigen composition they can be used for determination of particular species.



| Spot No. | Antigen | Description | <u>Pathogen</u> |
|----------|----------|--|--------------------------|
| A1 | МОМР Ср | Dominant major outer membrane protein (species specific) – structural protein; metabolic function | |
| A2 | MOMP1 | MOMP isoform, produced by posttranslational modification | |
| A3 | OMP2 Cp | Outer membrane protein (species specific) – structural protein of Chlamydia outer membrane complex | Chlamydia |
| A4 | OMP4 | Outer membrane protein | pneumoniae |
| A5 | OMP5 | Outer membrane protein | |
| A6 | P54 | Immunodominant outer antigen, highly specific to <i>Ch. pneumoniae</i> – sensitive marker for diagnosis of acute infection | |
| A7 | MOMP Ct | Dominant major outer membrane protein (species specific) – structural protein; metabolic function | |
| A8 | OMP2 Ct | Outer membrane protein (species specific) – structural protein of Chlamydia trachomatis Chlamydia outer membrane complex | Chlamydia trachomatis |
| A9 | HSP60 | Heat shock protein (GroEL); marker of chronic infection | |
| A10 | MOMP Cps | Dominant major outer membrane protein (species specific) – structural protein; metabolic function | Chlamydia |
| A11 | OMP2 Cps | Outer membrane protein (species specific) – structural protein of Chlamydia outer membrane complex | psittaci |

Test characteristics

Parameters of Microblot-Array Chlamydia IgA

| | Diagnostic Sensitivity | Diagnostic Specificity |
|-----------------|-------------------------------|-------------------------------|
| Ch. pneumoniae | 94.4% (n = 54) | 94.3% (n = 53) |
| Ch. trachomatis | 94.1% (n = 68) | 94.6% (n = 50) |

Parameters of Microblot-Array Chlamydia IgG

| | Diagnostic Sensitivity | Diagnostic Specificity |
|-----------------|-------------------------------|-------------------------------|
| Ch. pneumoniae | 94.6% (n = 111) | 96.0% (n = 25) |
| Ch. trachomatis | 98.3% (n = 41) | 92.7% (n = 60) |

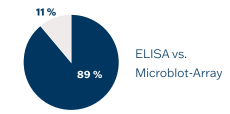
Comparative study

Correlation of results IgG

Ch. pneumoniae

| <u>n = 52</u> | Microblot-Array | ELISA | |
|---------------|-----------------|-------|--|
| positive | 31 | 32 | |
| negative | 15 | 20 | |

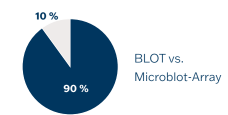
| total conformity | 88.5 % |
|------------------|--------|
| , | |



Ch. pneumoniae

| <u>n = 89</u> | Microblot-Array | BLOT |
|---------------|-----------------|------|
| positive | 73 | 81 |
| negative | 7 | 8 |
| | | • / |

| total conformity | 89.9 % |
|------------------|---------|
| total comorning | 0010 // |

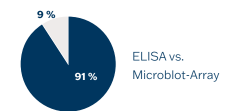


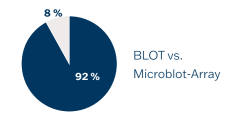
Ch. trachomatis

| <u>n = 70</u> | Microblot-Array | ELISA |
|------------------|------------------------|------------|
| positive | 17 | 20 |
| negative | 47 | 50 |
| total conformity | 91.4 | - % |

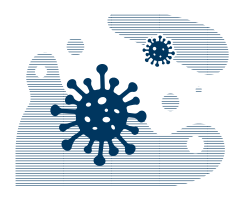


| <u>n = 39</u> | Microblot-Array | BLOT |
|------------------|-----------------|------|
| positive | 17 | 20 |
| negative | 19 | 19 |
| total conformity | 92.3 | % |



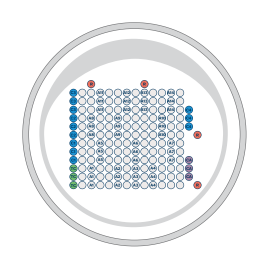






Microblot-Array for the diagnostics of SARS-CoV-2 and other coronaviruses

Microblot-Array COVID-19 kits enable simultaneous detection of multiple SARS-CoV-2 markers (NP, RBD, Spike S1, Spike S2, Spike S1 α -variant, Spike S1 γ -variant, Spike S1 δ -variant, E, ACE2, and PLPro). The kits also contain antigens to exclude cross-reactivities with other endemic coronaviruses (MERS-CoV, SARS-CoV, etc.). The kit contains antigens for the detection of various α , γ , δ mutations. The kits are optimized and validated for detection of IgA, IgG and IgM antibodies in human serum or plasma. They can be used for confirmatory testing, screening, epidemiological studies, identification of donors for convalescent plasma therapy, and other IVD and research applications related to the novel coronavirus.



| Spot No. | Antigen | Description | Pathogen |
|------------|-------------------------|--|------------|
| A1 | Nucleocapsid NP | A potent immunodominant coronavirus antigen that contains diagnostically important epitopes for the diagnosis of SARS-CoV-2 Sensitive detection of anti-SARS-CoV-2 IgG antibodies | |
| A2 | RBD | Receptor-binding domain of the S1 subunit of the spike (S) protein of SARS-CoV-2 Anti-RBD SARS-CoV-2 antibodies are highly subtype specific and protective The presence of anti-RBD antibodies significantly correlates with the formation of neutralizing antibodies IgA: for monitoring the immune response after a positive PCR reaction; indicator of the onset of the immune response IgM, IgG: detection of antibodies from 2 to 4 weeks after infection | |
| А3 | Spike S1 | The S1 subunit of the SARS-CoV-2 spike protein contains a receptor-binding domain (RBD), through which the virus binds to the surface of the host cell Anti-S1 antibodies are highly subtype specific, showing high sensitivity against SARS-CoV-2 and are protective | SARS-CoV-2 |
| A4 | Spike S2 | S2 subunit of the spike protein SARS-CoV-2 Plays an important role in the fusion of the virus with the cell membrane | |
| A 5 | Spike S1 α-variant | British mutation , Spike Glycoprotein S1 (B.1.1.7) | |
| A6 | Spike S1 γ-variant | Brazilian mutation, Spike Glycoprotein S1 (P.1) | |
| A7 | Spike S1 δ-varianta | Indian mutation, Spike Glycoprotein S1 (B1.617.2) | |
| A8 | Envelope protein (E) | The smallest major structural protein Important for different stages of viral infection and replication, important role in the life cycle of the virus | |

| Spot No. | Antigen | Description | Pathogen |
|----------|-----------------|--|-------------------|
| A9 | ACE2 | Angiotensin Converting Enzyme (transmembrane glycoprotein) A key component of the renin-angiotensin system Expressed in vascular endothelial cells in the heart, kidneys, but also the testes, liver, intestines, lungs and also the brain Involved in the regulation of cardiovascular and renal function | Human receptor |
| A10 | PLpro | Papain-like protease One of the basic SARS-CoV-2 proteins, essential for virus replication; deubiquitination activity Necessary for proteolysis of the viral polyprotein | SARS-CoV-2 |
| A11 | MERS-CoV S1 | Middle East Respiratory Syndrome Coronavirus S1 protein | |
| A12 | SARS-CoV Np | Severe Acute Respiratory Syndrome Coronavirus Nucleocapsid protein | Other endemic |
| A13 | HCoV 229E Np | Human coronavirus 229E Nucleocapsid protein | coronaviruses |
| A14 | HCoV NL63 Np | Human coronavirus NL63 Nucleocapsid protein | |

Test characteristics

Parameters of Microblot-Array COVID-19 kits

| | Diagnostic Sensitivity | Diagnostic Specificity |
|------------------------------|-------------------------------|-------------------------------|
| Microblot-Array COVID-19 IgA | 98.3% (n = 233) | 96.2% (n = 593) |
| Microblot-Array COVID-19 lgG | 98.7% (n = 309) | 99.3% (n = 600) |
| Microblot-Array COVID-19 IgM | 97.7% (n = 219) | 99.3% (n = 598) |

Comparative study

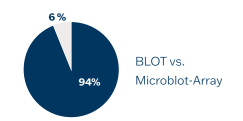
Correlation of results IgG

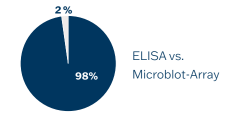
| n = 102 | Microblot-Array | BLOT |
|------------------|-----------------|------|
| positive | 87 | 91 |
| negative | 4 | 11 |
| total conformity | 93.5 | % |

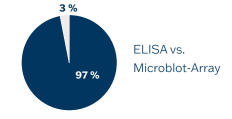
| n = 247 | Microblot-Array | ELISA |
|------------------|-----------------|-------|
| positive | 237 | 236 |
| negative | 10 | 7 |
| total conformity | 98.4 | % |

Correlation of results IgM

| <u>n = 228</u> | Microblot-Array | ELISA |
|------------------|------------------------|-------|
| positive | 193 | 193 |
| negative | 35 | 27 |
| total conformity | 96.5 | % |



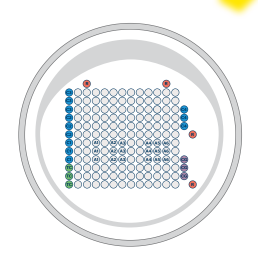






Microblot-Array for the diagnostics of cytomegalovirus infection

The Microblot-Array CMV kit is a new generation kit exhibiting a high diagnostic sensitivity and specificity and enabling quantitative evaluation. It is intended for confirmatory determination of specific antibodies in samples identified as positive or borderline by EIA or other serological tests. It is also intended to determine the presence of specific antibodies against CMV antigens, which allows to distinguish whether the primary infection is in an early or late stage or whether it is a secondary infection or reactivation.



Test characteristics

Parameters of Microblot-Array CMV kits

| | Diagnostic Sensitivity | Diagnostic Specificity |
|-------------------------|-------------------------------|-------------------------------|
| Microblot-Array CMV IgG | 98.1% | 99.9% |
| Microblot-Array CMV IgM | 96.9% | 99.1% |

| Spot No. | Antigen | Description |
|----------|--------------|--|
| A1 | p150 | Tegument protein UL32 A strong immunogen of the late stage of infection (late antigen); it does not develop in the early stage. Detectable in the IgG class in higher titres even in reactivation. |
| A2 | IEA (p72) | Immediate early antigen, capsid protein UL123 Plays a role in the early phase of the replication cycle of human CMV Important function in defence mechanisms against CMV infection |
| A3 | p65 | Tegument protein UL83 In the IgM class – one of the markers of the early stage of infection In the IgG class – rather typical for the late stage or infection reactivation |
| A4 | p52 | CM2 protein; UL44 In the IgM class – an important marker of the early stage of primary infection In the IgG class – reactivity rather in the late stage, or infection reactivation |
| A5 | p28 | Tegument protein UL99 A strong immunogen: it may develop in late stages of infection |
| A6 | gB | Membrane glycoprotein B Antibody response in IgG class – approximately 50–100 days after primary infection |

Interpretation of Microblot-Array CMV results

IgM IgG

| | p150 | IEA (p72) | p65 | p52 | p28 | gB | p150 | IEA (p72) | p65 | p52 | p28 | gB |
|--------------------------|------|--------------|-----|-----|-----|----|------|--------------|-----|-----|-----|-----|
| Early primary infection | - | (+) | + | + | - | - | _ | (+) | - | - | - | _ |
| Primary infection | (+) | (+) | + | +/- | (+) | - | - | (+) | (+) | (+) | (+) | - |
| Late primary infection | + | +/- | +/- | +/- | (+) | - | + | (+) | + | + | (+) | (+) |
| Persistence of infection | - | - | - | - | - | - | + | +/- | + | + | (+) | + |
| Reactivation | +/- | (+) | + | + | (+) | - | + | (+) | + | + | (+) | + |

(+) the marker may or may not be present

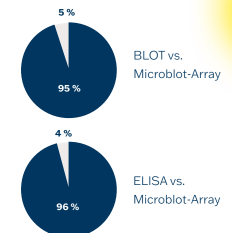
± weak or unclear reaction

Comparative study

Correlation of results IgG

| | Microblot-Array | BLOT |
|------------------|-----------------|------|
| positive | 31 | 31 |
| negative | 10 | 10 |
| total conformity | 9F 1 | 0/ |

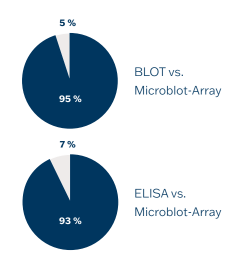
| | Microblot-Array | ELISA |
|------------------|-----------------|-------|
| positive | 200 | 199 |
| negative | 50 | 51 |
| total conformity | 96.4 | - % |



Correlation of results IgM

| | Microblot-Array | BLOT |
|------------------|------------------------|------|
| positive | 17 | 17 |
| negative | 21 | 21 |
| total conformity | 94.7 | % |

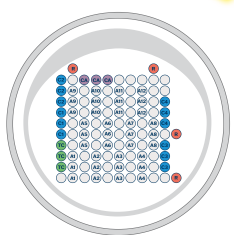
| | Microblot-Array | ELISA |
|------------------|-----------------|-------|
| positive | 109 | 110 |
| negative | 146 | 145 |
| total conformity | 93.3 | % |





Microblot-Array for the diagnostics of Epstein-Barr virus

Microblot-Array EBV kits are optimized and validated for detection of IgA, IgG and IgM antibodies in human serum or plasma. The kits are intended for confirmatory determination of specific antibodies in samples that have been identified mainly as positive or borderline by ELISA or other serological tests. Determination of specific class antibodies against EBV antigens is a useful tool for identifying a stage of EBV infection (primary infection, latent chronic infection or reactivation).



| Spot No. | Antigen | Description |
|----------|--------------|---|
| A1 | EBNA-1 | Epstein-Barr nuclear antigen 1 IgG: an important diagnostic marker of the late phase or reactivation of the infection IgM: the antibodies are detectable 2–4 months after primary EBV infection, they may also appear during reactivation |
| A2 | EBNA-2 | Epstein-Barr nuclear antigen 2 IgG: high antibody titres are present during chronic infection or in the post-acute phase The absence of IgG anti-EBNA-2 antibodies and the presence of anti-EBNA-1 antibodies rules out primary infection |
| А3 | VCA p18 | Viral Capsid Antigen p18; IgA: marker of primary infection; high titres persist in patients with nasopharyngeal carcinoma IgM: marker of primary infection; they may also be present during infection reactivation IgG: an important marker of the late phase of the infection, antibodies do not occur in primary infections |
| A4 | VCA p23 | Viral Capsid Antigen p23 Antibodies against this antigen can be detected during all phases of the infection (both IgG and IgM), they persist in the body for a long time |
| A5 | EA-D p54 | Early Antigen Diffuse p54; BMRF1 IgA: produced during primary infection; high titres during reactivation; high titres persist in patients with nasopharyngeal carcinoma An additional marker of acute EBV infection, detectable even in the latent phase of primary infection (both IgG and IgM) |
| A6 | EA-D p138 | Early Antigen Diffuse p138 IgA: produced during primary infection; high titres during reactivation; high titres persist in patients with nasopharyngeal carcinoma An additional marker of acute EBV infection, detectable even in the latent phase of primary infection (both IgG and IgM) |
| A7 | EA-R | Early Antigen Restricted protein p85; IgG: antibodies usually occur at a later stage; they are practically absent during the acute phase except in children; high levels in patients with reactivation or in immunocompromised patients |
| A8 | Rta | Replication and transcription Activator (BRLF1); A very early antigen IgG: a potential diagnostic marker of a nasopharyngeal carcinoma |

| Spot No. | Antigen | <u>Description</u> |
|----------|----------------|---|
| А9 | ZEBRA | Z Epstein-Barr replication activator protein; Trans-activator protein BZLF1 IgM: it is a very early indicator of an acute infection IgG: it is an early stage marker but it is also detectable during the late stages of the infection Serological marker of EBV reactivation, marker of EBV-associated diseases |
| A10 | gp85 | Probable membrane antigen gp85 (BDLF3); |
| A11 | gp350 | Epstein-Barr virus envelope glycoprotein gp350 (BLLF1); IgM: high titres in patients with infectious mononucleosis IgG: the titre increases only a few months after the primary infection Specific immune response for EBV-associated diseases |
| A12 | LMP1 | Latent membrane protein 1 Frequent in latent infections Linked to EBV-associated malignancies (nasopharyngeal carcinoma) |

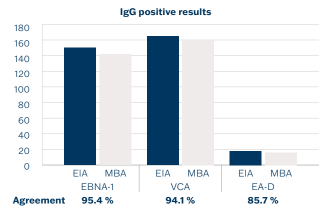
Test characteristics

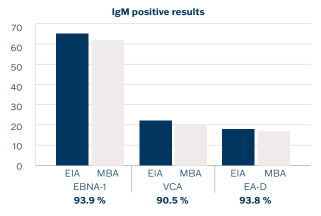
Parameters of Microblot-Array EBV kits

| | Diagnostic Sensitivity | Diagnostic Specificity |
|-------------------------|------------------------|------------------------|
| Microblot-Array EBV IgA | 98.9% (n = 167) | 96.7% (n = 70) |
| Microblot-Array EBV IgG | 98.8% (n = 167) | 96.9% (n = 70) |
| Microblot-Array EBV IgM | 96.4% (n = 61) | 89.3% (n = 60) |

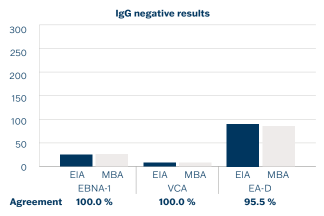
Comparative study

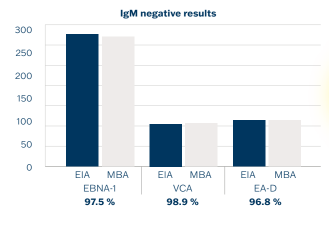
POSITIVE SAMPLES





NEGATIVE SAMPLES

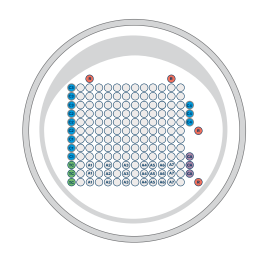






Microblot-Array for the diagnostics of *Helicobacter pylori* infection

The kits are optimized and validated for the detection of IgA and IgG antibodies against recombinant antigens *Helicobacter pylori* in human serum. For confirmation of ELISA positive or ambiguous results.



Test Characteristics

Parameters of Microblot-Array Helicobacter pylori kit

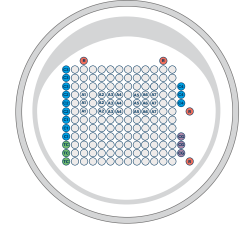
| | Diagnostic Sensitivity | Diagnostic Specificity |
|----------------------------------|-------------------------------|-------------------------------|
| Microblot-Array Helicobacter IgA | 96.5% | 99.1% |
| Microblot-Array Helicobacter IgG | 97.4% | 99.0% |

| Spot No. | <u>Antigen</u> | Description |
|----------|----------------|--|
| A1 | CagA, p120 | Cytotoxin associated gene A, highly specific, virulence factor |
| A2 | VacA, p87 | Vacuolating cytotoxin A, highly specific, virulence factor |
| А3 | UreA, p29 | Light subunit of urease, specific, virulence factor |
| A4 | NAP | Neutrophil-activating protein, virulence factor, potential biomarker of gastritis |
| A5 | НраА | Helicobacter pylori adhesin A, surface lipoprotein, potential biomarker of gastritis and gastric ulcer |
| A6 | НсрС | Helicobacter cystein-rich protein, virulence factor |
| A7 | GroEL | Chaperonin, heat shock protein (Hsp 60), virulence factor, considered as a marker of chronic infection |



Microblot-Array for the diagnostics of Herpes simplex virus infection

The Microblot-Array HSV kit is a new generation kit exhibiting a high diagnostic sensitivity and specificity and enabling quantitative evaluation. It is intended for confirmatory determination of specific antibodies in samples identified as positive or borderline by EIA or other serological tests. It is also intended to determine the presence of specific antibodies against HSV 1+2 antigens, which allows to distinguish whether the primary infection is in an early or late stage or whether it is a secondary infection or reactivation.



Test Characteristics

Parameters of Microblot-Array HSV kits

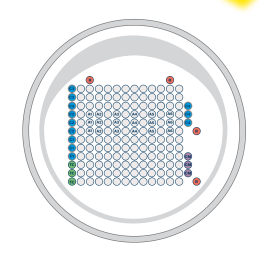
| | Diagnostic Sensitivity | Diagnostic Specificity |
|-----------------------------|-------------------------------|-------------------------------|
| Microblot-Array HSV 1+2 IgG | 99.9% | 97.5% |
| Microblot-Array HSV 1+2 IgM | 95.0% | 99.4% |

| Spot No. | Antigen | <u>Description</u> |
|----------|--------------|---|
| A1 | HSV 1+2 | Native HSV-1and HSV-2 antigen |
| A2 A3 | gC-1 gC-2 | Glycoprotein C-1 specific for Herpes simplex 1 virus; Glycoprotein C-2 specific for Herpes simplex 2 virus; Early antibody production |
| A4 A5 | gD-1 gD-2 | Glycoprotein D-1 specific for Herpes simplex 1 virus; Glycoprotein D-2 specific for Herpes simplex 2 virus serves to capture and entry of the virus into a potential host cell; stimulates high production of neutralizing antibodies, high similarity between HSV-1 and -2 |
| A6 A7 | gG-1 gG-2 | Glycoprotein G-1 specific for Herpes simplex 1 virus; Glycoprotein G-2 specific for Herpes simplex 2 virus Appropriate for differentiating between HSV-1 and -2 infection In the IgG class – indications of previous or probably latent infection; antibodies are formed only in the convalescent phase, they have been found also in patients with reactivation of infection In the IgM class – antibodies are produced only in the convales |



Microblot-array for the diagnostics of *Mycoplasma* infection

Microblot-Array Mycoplasma kits are used for the detection of specific IgA and IgG antibodies against recombinant antigens of *Mycoplasma pneumoniae* in human serum or plasma. Intended use is for confirmation of EIA ambiguous or positive results.



Test Characteristics

Parameters of Microblot-Array Mycoplasma pneumoniae

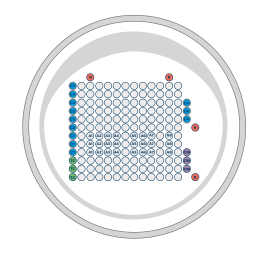
| | Diagnostic Sensitivity | Diagnostic Specificity |
|--------------------------------|-------------------------------|-------------------------------|
| Microblot-Array Mycoplasma IgA | 97.1% | 99.3% |
| Microblot-Array Mycoplasma IgG | 95.7% | 99.0% |
| Microblot-Array Mycoplasma IgM | 98.9% | 99.3% |

| Spot No. | Antigen | Description |
|----------|---------|---|
| A1 | P1 | Adhesin; the most important protein, a major virulence factor |
| A2 | p30 | Cytadhesin p30; the second most important protein, a major virulence factor |
| A3 | p116 | Adhesin, a major virulence factor |
| A4 | p65 | Surface protein; proline-rich P65 protein |
| A5 | HMW3 | Cytadherence high molecular weigh 3; adhesion-promoting protein |
| A6 | Mgp3 | Adhesion-promoting protein |



Microblot-Array for the diagnostics of *Yersinia* infection

The kits are suitable for the detailed determination of anti-Yersinia species specific IgA and IgG antibodies in human serum or plasma. Confirmation of ELISA positive or ambiguous results.



Test Characteristics

Parameters of Microblot-Array Yersinis sp.

| | Diagnostic Sensitivity | Diagnostic Specificity |
|------------------------------|-------------------------------|-------------------------------|
| Microblot-Array Yersinia IgA | 96.1% | 99.9% |
| Microblot-Array Yersinia IgG | 95.5% | 99.9% |

| Spot No. | Antigen | Description |
|----------|------------|--|
| A1 | YopB | Yersinia outer protein, transmembrane protein |
| A2 | YopD | Yersinia outer protein, transmembrane protein |
| A3 | YopM | Yersinia outer protein |
| A4 | YopN | Yersinia outer protein |
| A5 | LcrV | Low calcium response Virulence, important for YopD a YopB secretion |
| A6 | Ail | Attachment-invasion locus protein early phase, involved in the adhesion and invasion process, allows yersinia to survive outside the host cell, a significant virulence factor |
| A7 | Invasin | Surface adhesin binding to $\beta 1$ integrins on surface of target cells; important in the first stage of infection, a virulence factor |
| A8 | YscM-Y.Ent | Yop proteins translocation protein M |

Microblot-Array kits in development

Gastrointestinal diseases

| <u>Kit</u> | Antigens/parameters | <u>Availability</u> |
|---|---|---------------------|
| Microblot-Array Autoimmune gastroenteritis IgA, IgG | ASCA, DAG, tTG, IF, APCA, MPO | 2023 |
| Microblot-Array Gastro panel IgA, IgG | Helicobacter pylori, Yersinia enterocolitica, Autoimmune gastroenteritis | In development |

Herpetic infections

| Kit | Antigens/parameters | <u>Availability</u> |
|--|---------------------------|---------------------|
| Microblot-Array Herpetic infections panel IgG, IgM | EBV, CMV, HSV, VZV, HHV-6 | In development |

Liver-Kidney diseases

| Kit | <u>Parameters</u> | Availability |
|--------------------------------------|--|---------------------|
| Microblot-Array Liver-Kidney profile | 3E (BPO), M2, Sp100, PML, gp210, LKM-1, LC-1, SLA/LP, Ro52, ASGPR, Nup62, OGDC-E2, PDC-E2 | 2023 |

Atypical respirations

| <u>Kit</u> | <u>Parameters</u> | Availability |
|---------------------------------------|--|---------------------|
| Microblot-Array Atypical respirations | Mycoplasma pneumoniae, Bordetella sp., | In dovolopment |
| panel IgA, IgG, IgM | Chlamydia pneumoniae, Legionella pneumophila | In development |

Autoimmune neurological diseases

| <u>Kit</u> | Antigens | Availability |
|---|--|---------------------|
| Microblot-Array Paraneoplastic syndrome | Amphysin, CV2, GAD65, Hu, MA1, MA2, MAG, Recoverin, Ri, SOX1, Titin, Tr, Yo, ZIC4, AChR, MusK, Aquaporin-4 | In development |
| Microblot-Array Limbic encephalitis | LGI1, CASPR2, NMDAR, AMPA1/2, GABA1/2 | In development |

Anti-neutrophil cytoplasm antibodies

| <u>Kit</u> | <u>Antigens</u> | Availability |
|----------------------|-----------------|---------------------|
| Microblot-Array ANCA | PR3, MPO, GMB* | In development |

^{*}Other antigens will be added later.

Tropical diseases

| <u>Kit</u> | <u>Parameters</u> | Availability |
|-------------------------|---|---------------------|
| Microblot-Array | Dengue, Chikungunya, Zika, West Nile Fever, | In dayalanmant |
| Tropical diseases panel | Plasmodium, Rickettsia, Leptospira | In development |

Vector transmitted infections

| <u>Kit</u> | <u>Parameters</u> | Availability |
|------------------------------|--|---------------------|
| | Rickettsia, Babesia, Anaplasma phagocytophilum, | |
| Microblot-Array Vector | Neoehrlichia (Candidatus Neoehrlichia | In development |
| transmitted infections panel | mikurensis), TBEV, Borrelia burgdorferi, | in development |
| | Francisella tularensis, Q-fever (Coxiella burnetti)* | |

^{*}Kit composition will be more specified later

Endocrine antibodies

| <u>Kit</u> | Antigens/parameters | Availability |
|---|--|---------------------|
| Microblot-Array Thyreoid disease | TPO, TSH, TG | In development |
| Microblot-Array Diabetes mellitus Type I | ICA, IAA, IA-2, GAD, ZnT8 | In development |
| Microblot-Array Endocrine antibodies | Thyreoid disease, Diabetes mellitus Type I | In development |

TORCH

| <u>Kit</u> | <u>Parameters</u> | <u>Availability</u> |
|-----------------------------|-----------------------------------|---------------------|
| Microblot-Array TORCH panel | Toxoplasma, Rubella, CMV, HSV 1+2 | In development |

^{*} Expected availability may change, there may be a slight changes in antigenic composition. Status "In development" does not guarantee final launch of the product.

Ordering information



Kits

AUTOIMMUNITY

| Code | <u>Products</u> | No. of tests |
|----------|---------------------------|--------------|
| ANApMA96 | Microblot-Array ANA plus* | 96 |

^{*}Check availability in your country.

INFECTIOUS SEROLOGY

| Code | Products | No. of tests |
|----------|----------------------------------|--------------|
| BpAMA48 | Microblot-Array Bordetella IgA | 48 |
| BpGMA48 | Microblot-Array Bordetella IgG | 48 |
| BpMMA48 | Microblot-Array Bordetella IgM | 48 |
| BGMA096 | Microblot-Array Borrelia IgG | 96 |
| BMMA096 | Microblot-Array Borrelia IgM | 96 |
| CAMA096 | Microblot-Array Chlamydia IgA | 96 |
| CGMA096 | Microblot-Array Chlamydia IgG | 96 |
| CoVAMA96 | Microblot-Array COVID-19 IgA | 96 |
| CoVGMA96 | Microblot-Array COVID-19 lgG | 96 |
| CoVMMA96 | Microblot-Array COVID-19 IgM | 96 |
| CMGMA48 | Microblot-Array CMV IgG | 48 |
| CMMMA48 | Microblot-Array CMV IgM | 48 |
| EBAMA96 | Microblot-Array EBV IgA | 96 |
| EBGMA96 | Microblot-Array EBV IgG | 96 |
| EBMMA96 | Microblot-Array EBV IgM | 96 |
| НрАМА48 | Microblot-Array Helicobacter IgA | 48 |
| HpGMA48 | Microblot-Array Helicobacter IgG | 48 |
| HSGMA48 | Microblot-Array HSV 1+2 lgG | 48 |
| HSMMA48 | Microblot-Array HSV 1+2 IgM | 48 |
| MyAMA48 | Microblot-Array Mycoplasma IgA | 48 |
| MyGMA48 | Microblot-Array Mycoplasma IgG | 48 |
| MyMMA48 | Microblot-Array Mycoplasma IgM | 48 |

| Code | Products | No. of tests |
|---------|------------------------------|--------------|
| YAMA048 | Microblot-Array Yersinia IgA | 48 |
| YGMA048 | Microblot-Array Yersinia IgG | 48 |

Hardware a Software

| Code | Products |
|-----------|---|
| ARCXIX096 | Microblot-Array Reader (Array Reader C-series) + Software |

Components

| Code | Products |
|-----------|------------------------------|
| 000008262 | Universal Solution (300 ml)* |

^{*}In the case of automated processing, an additional universal solution is required because of the dead volumes of the instruments. We recommend 2 extra bottles/kit (when running one plate per week). Please contact our sales representatives for more information.





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Company is certified to the quality management system standards ISO 9001 and ISO 13485 for in vitro diagnostics.