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REVIEW

Challenges in Detection of Serum Oncoprotein: Relevance to Breast Cancer Diagnostics

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Received: 30 July 2021 Accepted: 2 October 2021 Published: 14 October 2021 Abstract: Breast cancer is a highly prevalent malignancy that shows improved outcomes with earlier diagnosis. Current screening and monitoring methods have improved survival rates, but the limitations of these approaches have led to the investigation of biomarker evaluation to improve early diagnosis and treatment monitoring. The enzyme-linked immunosorbent assay (ELISA) is a specific and robust technique ideally suited for the quantification of protein biomarkers from blood or its constituents. The continued clinical relevancy of this assay format will require overcoming specific technical challenges, including the ultrasensitive detection of trace biomarkers and the circumventing of potential assay interference due to the expanding use of monoclonal antibody (mAb) therapeutics. Approaches to increasing the sensitivity of ELISA have been numerous and include employing more sensitive substrates, combining ELISA with the polymerase chain reaction (PCR), and incorporating nanoparticles as shuttles for detection antibodies and enzymes. These modifications have resulted in substantial boosts in the ability to detect extremely low levels of protein biomarkers, with some systems reliably detecting antigen at sub-femtomolar concentrations. Extensive utilization of mAb therapies in oncology has presented an additional contemporary challenge for ELISA, particularly when both therapeutic and assay antibodies target the same protein antigen. Resolution of issues such as epitope overlap and steric hindrance requires a rational approach to the design of diagnostic antibodies that takes advantage of modern antibody generation pipelines, epitope binning techniques and computational methods to strategically target biomarker epitopes. This review discusses technical strategies in ELISA implemented to date and their feasibility to address current constraints on sensitivity and problems with interference in the clinical setting. The impact of these recent advancements will depend upon their transformation from research laboratory protocols into facile, reliable detection systems that can ideally be replicated in point-of-care devices to maximize utilization and transform both the diagnostic and therapeutic monitoring landscape.

Keywords: breast cancer, diagnosis, ELISA, detection, plasmonics, sensitivity

Introduction

Breast cancer is the second most prevalent type of cancer to afflict women, behind only cancers of the skin. The American Cancer Society estimates that 268,000 women were diagnosed with invasive breast cancer in 2019 in the United States, along with 3.8 million survivors with a previous diagnosis.¹ While prognoses vary among breast cancer subtypes, it has been well established that patient outcomes are significantly improved with earlier diagnosis. Current 5-year survival rates for localized and regional breast cancer at the time of diagnosis are 99% and 86%,

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respectively, which drops precipitously to 28% when a patient is diagnosed late-stage with distant metastases.² Screening methods such as mammography and ultrasonography have increased the early detection of breast cancers, but the limitations of these methods^{3,4} have driven efforts to identify biomarkers that could be utilized for early-stage diagnosis as well as evaluation of treatment efficacy.

The most desirable method for diagnosing and monitoring therapeutic effectiveness is to measure key molecular characteristics in biofluids, specifically blood and its constituents. An immunoassay of protein biomarkers in biological samples by enzyme-linked immunosorbent assay (ELISA) is ideally suited for this purpose. ELISA is a highly specific, minimally invasive, and cost-effective technique to better understand the pathological and biological status of a patient. ELISA has long been the gold standard immunoassay for quantifying proteins in biofluid, and the ever-expanding list of clinically relevant biomarkers for disease diagnosis and treatment management drives the need for next-generation assays that are specific and sensitive enough to accurately quantify trace amounts of protein. While ELISA is a specific, sensitive, and reliable assay methodology, significant technical challenges have evolved with its modern clinical application.

As proteins are the main drivers of biological function and phenotype, the evaluation of pertinent protein biomarkers can provide the most relevant patient status assessment regarding the diagnosis of disease or evaluation of therapeutic success. Targeted protein quantification from biofluids for biomarkers of interest offers a snapshot in time of a patient's health status, as the molecular profile of blood is directly associated with their physiological state. ELISA analysis has historically been applied to a large collection of pathologies, ranging from the identification of blood-borne infectious agents to the detection of markers for various cancers and neurological diseases.^{5–8} This is a challenging pursuit for a single assay format, as the broad range of clinically relevant concentrations of disease-related biomarkers in blood can range from the pg mL^{-1} (eg, Interleukin-6 in oral cancer) to the ng m L^{-1} range as is the case for the extracellular domain (ECD) of the human epidermal growth factor receptor-2 (HER2) in breast cancer.⁷ The limit of detection for a traditional colorimetric ELISA however, which is dependent on several factors, can range from 0.01 to 1.5 ng mL^{-1,9,10} highlighting the need for more sensitive assays.

Many iterations of the traditional ELISA have been developed that significantly improve the sensitivity of the assay and continue to maintain its clinical relevancy. Because protein targets cannot be amplified to enhance their signal like nucleic acids, alternative signal-based amplification strategies have been developed to improve the sensitivity of ELISA. Approaches include utilizing more sensitive nanoparticle-based substrates,¹¹ combining ELISA with the polymerase chain reaction (PCR),¹² and incorporating nanoparticles as shuttles for detection antibodies and enzymes.^{13,14} These modifications have resulted in substantial boosts in the ability to detect extremely low levels of protein biomarkers, with some systems reliably detecting antigen at sub-femtomolar concentrations.¹⁵

The explosion of monoclonal antibody (mAb) therapies, currently the largest class of biopharmaceuticals, has presented an additional contemporary challenge for ELISA. Biomarker monitoring for evaluation of treatment efficacy is a powerful concept but is potentially impeded by the existence of therapeutic antibodies that may be present in biofluid samples. Particularly in oncology, where targeted antibody therapies have been increasingly employed, the accurate quantification of target proteins can be complicated by interfering therapeutic antibodies. Resolution of issues such as epitope overlap and steric hindrance requires a rational approach to the design of diagnostic antibodies that takes advantage of modern antibody generation pipelines, epitope binning techniques and computational methods to strategically target biomarker epitopes.¹⁶

This review covers strategies implemented to date that address two key technical areas crucial to the exact quantification of protein targets in biofluids with ELISA in the modern oncology era: amplification of signal to improve assay sensitivity and rational design of assay antibodies and their fragments to circumvent steric issues present when an antibody therapy and diagnostic assay target the same protein. Implementation of these strategies will facilitate improved clinical outcomes through earlier diagnosis and more accurate therapeutic monitoring.

Traditional ELISA

ELISA, a biochemical immunoassay that takes advantage of the specificity of antibody–antigen interactions to capture and detect antigen on a solid substrate, was first described in 1971.¹⁷ Variations of the assay have resulted in several different commonly employed methods to capture and detect antigen targets: Direct, Indirect, Competitive and Sandwich ELISA as represented in Figure 1.



Figure I Diagrammatic representation of different types of ELISA.

Direct and Indirect ELISA rely on direct adsorption of antigen to the plate surface for immobilization of target proteins from biological samples. The captured antigen of interest is specifically targeted for detection with either an enzyme-labeled primary antibody (Direct ELISA) or by employing an unlabeled primary antibody matched to an enzyme-labeled secondary antibody (Indirect ELISA). Competitive ELISA determines antigen concentration by evaluating the interference a patient sample has on an expected signal. Plate wells are coated with reference antigen, while test samples are incubated with labeled antibody specific to the antigen of interest. Resulting antibody/antigen complexes in the sample are thus unable to bind the reference antigen upon addition to plate wells, resulting in reduced signal compared to control wells lacking test sample.^{17,18}

Direct, Indirect, and Competitive ELISA variants utilize a sole antibody specific to the protein target and thus rarely reach the specificity of the Sandwich ELISA, which employs primary and secondary antibodies that are reactive to the protein of interest. This increased specificity makes the Sandwich variant the preferred diagnostic ELISA format. In this format, the primary antibody is adsorbed to the surface of the test plate well and used to capture antigen, which is then recognized by a secondary enzyme-labeled detection antibody that is targeted to a different epitope on the captured protein.¹⁸ Table 1 describes the advantages and disadvantages of different ELISA formats.

A key element of ELISA sensitivity relates to the affinity of the assay antibody for its antigen target, particularly the capture antibody in the Sandwich ELISA.^{9,10} Both mAb and polyclonal antibodies (pAb) have been applied to ELISA and fully optimizing the Sandwich ELISA format may result in the employment of both. Over the last 20 years, mAbs, which target a single epitope on an antigen, have come to the forefront of diagnostic testing due to their high specificity and affinity.¹⁹ Selection of host species for generating diagnostic antibodies is also a crucial decision for diagnostic assays. While the mouse was often the earliest host for producing ELISA antibodies, rabbits have become increasingly utilized due to several advantages over mice, including affinities up to 100 times greater, superior specificity, and higher stability.^{20–22} The rabbit immune response to foreign antigen also has a broader repertoire, enabling the recognition of epitopes that are not immunogenic in mice, especially in small molecules and peptide antigens.²³

Signal amplification methods are required to detect and quantify the bound protein of interest. This has traditionally been achieved by conjugating one of several different detection enzymes to primary (Direct ELISA) or secondary antibody (Indirect or Sandwich ELISA), most commonly horse-radish peroxidase (HRP) or alkaline phosphatase (AP), to generate signal through catalysis of substrate added in the terminal steps of the ELISA procedure.²⁴ HRP, an oxidoreductase that catalyzes the oxidation of ELISA substrates by hydrogen peroxide, is generally favored over AP for this role due to its higher stability and smaller size (40kDa vs 86kDa),²⁵ which enables more enzyme conjugates per detection antibody and thus more potent signal amplification.²⁴ The increased size of AP can also cause steric hindrance issues due to the proximity of the antigen-antibody complex, potentially resulting in lower enzyme activity.

A multitude of substrates produce different forms of quantifiable signal after catalysis by HRP or AP, with the most

Type of ELISA	Advantages	Disadvantages	
Direct	 Rapid and easy set-up Simple procedure 	 Less specific since it employs single antibody. Detection antibody should be conjugated. Potential for high background signal. 	
Indirect	 Amplification of signal using a secondary antibody. High sensitivity as compared to direct ELISA. Cost effective 	 Longer protocol due to greater number of steps. Potential for cross-reactivity caused by secondary antibody leading to non-specific signal. 	
Competitive	 Ability to quantitate small molecules with low analyte concentration. Sample purification not required prior to use. 	• Low sensitivity and specificity	
Sandwich	• Highest specificity and sensitivity Antigen purification not required prior to use.	 Costly and time-consuming. Requires matched antibody pairs. Long procedure. 	

Table I Advantages and Disadvantages of Different Types of ELISA

well-established substrate categories including, colorimetric, fluorescent, and chemiluminescent. Colorimetric substrates are the most leveraged and have several advantages in ELISA including low cost, simplicity, and a signal that is visible to the naked eye.²⁶ Fluorescent substrate catalysis results in the generation of a product that fluoresces when excited by light at the appropriate wavelength. While fluorescent substrates are marginally more sensitive than colorimetric, the dynamic range of the assay is also wider, allowing higher optical density (OD) measurement than the 4.0 OD limit imposed by most colorimetric plate readers. Chemiluminescent substrates produce photons of light when catalyzed by either HRP or AP, and can reach levels of sensitivity several orders of magnitude higher than fluorescent substrates.²⁶

Interference in ELISA

Biological samples analyzed by immunoassay are complex matrices that may contain components that interfere with the accurate quantification of a target analyte. In the clinic, interfering elements can be endogenous and inherent to the patient, or exogenous and due to the administration of a substance or therapy. Excessive levels of certain serum components, namely lipids and bilirubin, are the most reported causes of endogenous interference.^{27,28} Lipemia, or excessive levels of triglycerides in the blood, is usually visible to the naked eye and can be caused by recent dietary fat consumption or abnormal lipid metabolism and interferes due to light scattering or absorbance during signal quantification at the end of the ELISA procedure.²⁷

Icterus or jaundice is a condition in which blood concentrations of bilirubin are elevated, which can cause spectrophotometric interference due to bilirubin's strong absorbance of light between 340 and 500nm.²⁹ This absorbance range overlaps with the absorbance of products from ELISA substrate catalysis, obfuscating the true signal.³⁰ Caution must also be used in interpreting ELISA results from patients with rheumatoid arthritis, as this condition results in the generation of autoantibodies against the fragment crystallizable (Fc) portion of Immunoglobulin G (IgG) class of antibodies potentially skewing assay results through non-specific interaction with ELISA antibodies.^{31,32}

Regardless of the source, the distorting of assay results can occur via direct or indirect mechanisms. Bias through direct interference is induced when the interferent is quantified instead of the target analyte, through cross-reaction with antibody components of the testing assay. Indirect interference manifests when components interact with either a reagent or the target analyte, altering the affinity of the analyte for assay antibodies. The result of these offtarget effects can be either false-positive (an increase) or false-negative (a decrease) in the quantified signal, potentially leading to misdiagnosis or a misleading treatment evaluation.

Diagnosing ELISA interference can be achieved through spike-and-recovery experiments and analysis of serially diluted samples. Spiking biological samples with a known amount of target analyte enables the quantification of interference, as an assay free of interference should

accurately quantify the total amount of analyte added. Frequently, interfering components will dilute out of a sample, which will be apparent when the dilutionadjusted concentration is calculated from the same sample across a range of dilutions. Testing for parallelism, the ability of an assay to correctly measure an analyte across a dilutional range, can implicate serum matrix interference when analyte measurements display non-parallelism at low dilution which disappears as dilution increases. Implacable non-parallelism at higher dilutions likely indicates either a high concentration of interferent that dilution cannot remedy or an exceptionally high affinity of interferent for the analyte or assay components. In cases of high-affinity interferents such as antibodies, several approaches can be utilized to overcome or minimize the interference. Such techniques include acid-dissociation, which can disrupt the binding of the analyte to interferent.³³ choosing an optimal blocking agent or buffer,^{34,35} or targeted design of immunoassay antibodies that bind to free epitopes on the analyte in the presence of the impeding agent.

Interference through both endogenous and exogenous antibodies has been documented. Endogenous interference can occur in patients who have been exposed to murine antibodies through chance environmental exposure or deliberate antibody therapy, resulting in the production of human anti-mouse antibodies (HAMA) that potentially interfere in immunoassays built on mouse antibodies.³⁶ With the continuing growth of mAb therapies, which display high affinity for their targets, exogenous interference poses an increasing challenge for immunoassays, particularly for therapy-monitoring immunoassays used to analyze patient samples likely to contain high concentrations of these therapeutics. False-positive interference was reported with early antibody therapy for treating immunodeficient states.^{37–41} This likely under-reported phenomenon has the potential to skew immunoassay results and negatively affect patient outcomes that rely on ELISA for treatment guidance, highlighting the ongoing need for strategies that ensure the validity of results in diagnostic and therapy-tracking immunoassays.

Domain-Specific Targeting for ELISA Antibodies to Prevent Interference

Serum biomarker monitoring by ELISA to evaluate targeted therapy effectiveness is frequently reliant on quantifying the same protein that the mAb therapy targets.⁴² For instance,

in breast cancer, human epidermal growth factor receptor-2 (HER2; also known as ErbB2), a transmembrane tyrosine kinase receptor that is overexpressed in ~20% of breast cancer patients, drives a pro-oncogenic program of unchecked cell growth and apoptosis resistance associated with poor prognosis.43 The HER2 extracellular domain (ECD), the target antigen for current anti-HER2 mAb therapies such as trastuzumab and pertuzumab, is comprised of four subdomains (I-IV) and can be cleaved from the surface of breast cancer cells by extracellular-matrix degrading matrix metalloproteases (MMPs).44-46 A growing body of literature has demonstrated that a reduction in HER2 ECD concentrations in patient serum during therapy is predictive of pathologic complete response (pCR) and improved outcomes^{7,47-51} while rising or refractory serum HER2 ECD levels indicate resistance to therapy and the need for alternative treatments.³⁵ While the therapeutic binding domains of trastuzumab and pertuzumab have been characterized as within subdomain IV and subdomain II (Figure 2),⁵² respectively, commercially available ELISAs utilize antibodies that often bind to undefined or undisclosed epitopes on the HER2 ECD. This is potentially problematic, as inherent in therapeutic monitoring programs is the fact that the patient will have the mAb therapeutic in their serum during ELISA testing, potentially obfuscating the true level of HER2 ECD present.

When mAb therapy and immunoassays share an antigenic target, two mechanisms may bias attempts at quantification. The therapeutic antibody and immunoassay antibodies may have direct epitope overlap, binding to the same antigenic amino acid sequences, resulting in competition for the same physical location. Sharing of epitopes is not necessary for therapeutic antibodies to interfere, however, as antibody bound to neighboring epitopes can also provoke steric hindrance that physically blocks access of assay antibodies to their target epitopes. Overcoming these issues requires mapping of therapeutic mAb antigen binding sites as well as rational immunoassay antibody approaches that produce high affinity, epitope-specific antibodies to targeted domains that remain unbound in the presence of mAb therapeutics.

Various methods have been employed to generate mAbs for both therapeutic and immunoassay applications, including hybridoma and phage display technology. Hybridoma technology has historically been the predominant method of isolating mAbs and is a favored method due to the in vivo generation of highly specific antibodies that have undergone the natural affinity maturation



Figure 2 Three-dimensional structure of Her2-antibody complexes. (Left) Structure of the extracellular domain of Her2 is shown in ribbon representation. Subdomains I–IV are shown in pale-green, orange, green and purple colors, respectively. (Middle) Structure of Her2-Trastuzumab complex is shown. Trastuzumab (blue) binds to Sbd-IV. (Right) Structure of Her2-Pertuzumab complex is shown. Pertuzumab (pink) binds to Sbd-II (PDB: 1N8Z¹⁵⁹ and 1S78¹⁶⁰).

process. Immunization of the host animal, predominantly mice or rabbits, with antigen provokes the production of antibody in splenic plasma cells. Extracted spleen cells are fused with multiple myeloma-derived cells forming immortalized hybridoma cells that are sorted through dilution to produce cultures generating antibodies against a sole epitope.

Guiding this system to produce mAb against a specific domain can begin with immunizing the host animal against a truncated peptide fragment from a region of interest in the full-size antigen. However, this approach can have limited success⁴⁵ due to the variable nature of epitopes. Epitopes can be sequential and correspond directly to the antigen peptide sequence, or conformational and result from disparate residues brought into proximity in the folded protein that is not represented by linear peptide fragments. Furthermore, cysteine-rich proteins, like the HER2 ECD, form specific intramolecular disulphide bonds that mediate proper folding of the full-length protein and are difficult to recapitulate when recombinantly expressed in truncated form in vitro. For example, Rockberg et al immunized rabbits with recombinant proteins that were restricted to one of the four subdomains of the HER2 ECD to generate antibodies against individual subdomains.⁴⁵ Immunogens that contained entire isolated subdomains did not generate antibodies that recognized the native full-length protein, while an immunogen that overlapped subdomains II and III, that the authors speculate contained linear epitopes, produced functional antibody against native HER2 ECD.

The immunogenicity of a peptide fragment is not evenly distributed, as hydrophilic and N- or C-terminal residues are inherently more immunogenic than hydrophobic and central amino acid residues.^{53,54} This uneven distribution can produce antibodies to epitopes biased to the terminal ends of peptide fragments that may not exist in the full-length antigen. Strategies to target the immune response to specific epitopes on a truncated immunogen have been developed based on observations made about cell surface receptor glycosylation. Work by Peiris et al and others have demonstrated that the HER2 ECD is variably glycosylated depending on oncogenic status and epitope masking due to heavy glycosylation on the ECD reduces the binding and effectiveness of trastuzumab.55 Applications of this concept in recent reports demonstrated that masking the terminal amino acids of peptide antigens with weakly immunogenic glycosylation can shift the immune response toward central residues, providing a path toward the generation of effective domain-specific mAbs using peptide fragments.^{56,57}

Screening of immunized animal antiserum or hybridoma cell cultures for antibodies produced from full-length antigen can also isolate mAb against a targeted domain. Production of domain-specific recombinant forms of antigen by *Escherichia coli* or other systems, through the expression of plasmids with the antigenic gene restricted

to individual domains, can serve as controls to define mAb binding sites. Western blot analysis of these individual domains against candidate mAbs can be rapidly performed and verification against the non-denatured domain-specific antigen can be confirmed by immunocytochemistry in transfected cell lines expressing domain-restricted forms of the target antigen.^{58,59} More high-throughput, systematic methods of epitope mapping have classically utilized overlapping peptide arrays that enable sequential epitope mapping.^{60–63} Conformational epitopes, however, require screening with the full three-dimensional structure of the intact antigen. The site-directed epitope masking technique, which utilizes mutants of the full-size targeted antigen tethered to a solid phase, effectively blocks the binding of the antibody to the corresponding surface of the antigen.⁶⁴ Antibodies that can still access their corresponding epitopes and bind are subsequently depleted from serum, allowing the enrichment of antibodies reactive to the masked surface and identification of their binding sites. This technique was successfully used to generate A21, a mAb against HER2 ECD that exhibited proliferation inhibition activity on HER2 overexpressing cancer cell lines and has been effectively employed in ELISAs to quantify HER2 ECD.⁶⁵

In the context of an interfering mAb therapy bound to an ELISA target antigen, surface plasmon resonance (SPR) serves as a tool for selecting immunoassay mAbs that react to epitopes free of therapeutic antibody interference. SPR is an optical sensing technology in which antigen is immobilized on the surface of a probe to which different mAbs are serially exposed through microfluidic channels. The binding of the antibody to the antigen causes a shift in SPR signal and if subsequent antibodies react to the same epitope, they will be unable to bind the antigen and there will be no subsequent shift in SPR signal. This enables the sorting and grouping, or binning, of candidate assay antibodies based on whether they compete for the same epitope on their target antigen. Agnolon et al employed SPR to characterize and epitope bin antibodies used in an ELISA sandwich assay for the quantification of serum levels of HER2 ECD.⁶⁶ SPR was used to verify that both assay antibodies do not share epitopes with trastuzumab by exposing probe bound HER2 ECD to trastuzumab prior to the assay antibodies' exposure, confirming that all three can simultaneously bind their epitopes. These results were also validated with displacement testing, in which HER2 ECD was pre-incubated with trastuzumab at two concentrations (50 and 200ug/mL) prior to running the sandwich ELISA assay and compared to trastuzumab-naïve samples. Interference due to trastuzumab was minimal, with a decrease in the signal of 7.9%, and interestingly was independent of drug concentration suggesting an interference mechanism driven by steric hindrance, as opposed to competition for overlapping epitopes. As mAb therapies are increasingly employed, the need for simple and economical characterization of assay antibodies that SPR enables is essential to ensure accurate biomarker quantification.

While hybridoma technology has generated a plethora of assay and therapeutic antibodies used today, in vitro techniques based on recombinant DNA technology have rapidly become the fastest and most economical targeted antibody development method. Phage display technology, the most common in vitro technique for producing antibody, facilitates the screening of large libraries of antibodies or antibody fragments against a target antigen by forcing their expression on the surface of bacteriophages. Phage display libraries containing a pool of antibody genes are constructed by extracting mRNA from B-cells isolated from animals that have been immunized with the antigen of interest. Integration of B-cell antibody genes with a phage coat gene results in the display of fusion antibodycoat proteins on the surface of the phage, enabling characterization through repeated rounds of panning against the antigen of interest.⁶⁷ Phages that display antibody that binds the target antigen with high affinity adhere to the solid phase-bound antigen and are then eluted for sequencing and identification. While this is a rapid method to screen large numbers of antibodies, the panning process identifies only the highest affinity antibodies, limiting diversity and biasing "hits" away from potential epitopes or domains of interest with lower affinity to the target antigen. One recent approach to solving this issue involves the incorporation of noncanonical amino acids (ncAAs) into epitopes of interest in the antigen used for panning phage display libraries.⁶⁸ The advantage that ncAAs such *p*-benzoyl-L-phenylalanine and as p-azido-L-phenylalanine confer is their propensity to covalently cross-link proteins in the vicinity when exposed to UV light. Upon panning of phage, the photocrosslinker that forms between the epitope-specific ncAA and reactive antibody on the surface of phage after UV exposure enables the selection of antibodies specific to the target epitope containing the ncAA, regardless of affinity. This promising method allows for the creation of phage libraries from full-size antigen, thus preserving

conformational epitopes, while still enabling the selection of epitope-specific mAbs.

Phage display libraries can also be vehicles for displaying peptide fragment libraries to epitope map an existing antibody. The mAb A21, generated against HER2 ECD⁶⁵ was partially epitope mapped to subdomain I of the HER2 ECD by bio panning a random 12mer peptide fragment phage library against A21. Peptides that bound A21 were then isolated and sequenced for alignment with the full-length HER2 molecule to identify the location of the epitope.⁶⁹ Further resolution on the exact amino acids that comprise the A21 epitope was obtained by evaluating A21 binding to individual recombinant subdomains of the ECD in ELISA format, which resulted in a candidate epitope site in subdomain I. Subsequent mutagenesis analysis within subdomain I revealed several surface-exposed residues that were most impactful to the affinity of A21 for HER2 ECD. Characterization of ELISA mAb epitopes to HER2 ECD is vital to ensure non-overlap with mAb therapies and accurate quantification, but is also essential for anti-HER2 mAb therapeutics in particular, as has been demonstrated by the large body of data confirming that downstream cellular effects of mAb therapeutics against the HER2 ECD on cancer cells are variable depending on epitope location.^{70–75}

Computational methods have also been utilized to generate complementarity-determining design and regions (CDRs) of antibodies against specific domains of antigen targets, especially against weakly immunogenic epitopes. Historically, this method has relied on the three-dimensional crystal structure of the antibody, the antigen and the antibody-antigen complex to then model and design antibodies complementary to targeted epitopes.^{76–78} The success of these computationally designed antibodies relies heavily on precise models for accurate in silico optimization and continues to be a challenge,⁷⁹ though recent advances have been made in de novo design of antibody without an initial model structure. Specific de novo approaches vary and can involve epitope prediction based on specific antibodies.⁸⁰⁻⁸² and simulating the gene recombination process within B-cells to generate novel CDRs.^{83,84} These de novo approaches have continuously improved in producing antibodies with high affinity and specificity and, given the difficulty in accurately modeling the structure of antigen-antibody complexes, are likely to be the main driver in expanding the ability of computational methods to design future epitopetargeted assay antibodies.

Antibody Fragments in ELISA

Designing ELISA antibodies to circumvent interfering therapeutics can ultimately be successful. However, due to their large molecular weight (~150 kDa) limiting access to target epitopes, the physical constraints of multiple antibodies binding to the same antigen can reduce the efficacy of assay mAbs. Fortunately, the modular nature of antibodies has enabled the creation of antibody fragments with significantly reduced size which aids in accessing cryptic epitopes and reduces interference from select serum matrix components. These fragments are also more easily genetically manipulated and expressed in bacterial systems for faster cultivation and higher yields.^{85–88}

The gross structure of IgG, the most frequently utilized class of antibody for immunoassays, can be subdivided into the fragment antigen-binding (Fab) and fragment crystallizable (Fc) portions. The Fab region contains the constant and variable domain of the heavy and light chain of the antibody and is responsible for binding antigen, while the Fc portion is located on the tail section and interacts with Fc receptors on the surface of cells to modulate the activation of the immune system. The Fab fragment is the most established antibody fragment in both diagnostics and therapeutics^{89,90} and can be generated either recombinantly or enzymatically by utilizing papain to cleave IgG into two monovalent Fab fragments and a single Fc fragment. Besides this, reduced size with increased avidity over the single Fab fragment can be achieved by digesting IgG with pepsin, which cleaves below the IgG hinge region and produces a single bivalent F(ab')2 fragment, retaining both paratopes of the full-size IgG.⁹¹ Fab and F(ab')2 fragments retain the specificity of the full-size antibody while reducing the size to ~50 kDa and 110 kDa, respectively, potentially enhancing accessibility to partially blocked epitopes and allowing deeper penetration into tissue when utilized for immunohistochemistry. Furthermore, the employment of Fab/ F(ab')2 fragments eliminates the risk of assay interference due to rheumatoid factors, which bind the Fc portion of IgG.⁹²

Single chain variable fragments (scFv) are the smallest fragment of IgG that retains a complete antigen binding domain and are composed of a variable region from the heavy chain (VH) and light chain (VL) connected by a flexible synthetic linker molecule, effectively reducing its size to ~ 27 kDa. This small size makes them ideal as

screening candidates for phage display as well as largescale production in microbial systems.⁹³ While ELISA assays have been constructed from scFv fragments,⁹⁴ drawbacks to this format include decreased thermostability as well as dimerization and fragment aggregation due to the synthetic linker molecule.⁹⁵ These shortcomings have driven the search for additional antibody fragments similarly reduced in size that are not reliant on a synthetic component.

Further size reduction of ELISA antibody components can be achieved by utilizing an atypical class of antibodies found to be naturally occurring in camelids and several different types of sharks. These so-called heavy chain antibodies (HCAbs) lack the light chain variable domain typically found in IgG antibodies, with specificity for antigenic targets determined solely through the heavy chain variable domain. The cleavage of HCAbs to isolate the variable domain yields a fragment known as a single domain antibody (sdAb). Generation of sdAb from camelid antibody generates VHH (Variable domain of the Heavy chain of HCAbs) fragments, or nanobodies, which have a molecular weight of just 15kDa. HCAbs in sharks, termed immunoglobulin new antigen receptors (IgNARs), and the isolated variable domain, termed variable new antigen receptors (vNARs), are the smallest known antibody fragment at ~12kDA. Both VHH and vNARs possess advantages in solubility and stability over equivalent VH domains from human IgG.^{96,97} Each also has a uniquely long complementarity-determining region 3 (CDR3) that confers a structural advantage, enabling the binding of antigen cleft regions that are not as easily accessed by full size IgG.^{98,99} The small size and ability to access cryptic epitopes make VHH and vNARs ideal candidates for ELISA, especially in the context of circumventing an interfering component, such as a mAb therapeutic contained in a patient sample. Camelid VHH, in particular, have been employed to build ELISAs against antigens across a wide array of research fields, including oncology (HER2 extracellular domain), virology (HIV, rotavirus),^{100,101} and toxicology (ricin, cholera).102

The large body of work on antibody structure and function has provided ELISA designers a plethora of antibody fragments to choose from for assay optimization. These fragments offer great promise in solving the multiple issues inherent to biomarker monitoring. However, careful vetting of candidate antibody fragments is essential, as the advantages of smaller size for epitope access and easier production can be outweighed by the increased risk of aggregation and decreased thermal stability.

Signal Amplification

While antibody affinity and specificity are major limiting factors in ELISA, the accurate detection of bound antibody is also a critical determinant in assay sensitivity. Unlike systems that increase signal through target-based amplification methods, as PCR does with DNA, ELISA relies on signal-based amplification to catalytically increase the signal generated from a single binding event between assay antibody and target antigen. The colorimetric ELISA is the most common method employed in the clinical setting, due to its simplicity and low cost. However, this direct approach is offset by lower sensitivity for trace analytes compared with fluorescent and chemiluminescent platforms. Novel methods of signal amplification have been developed to increase ELISA sensitivity, across all signal transduction platforms, to address the clinical challenge of detecting minute concentrations of relevant biomarkers for diagnosing early-stage disease and therapeutic monitoring.

The most utilized method of ELISA signal amplification is the enzyme-mediated process catalyzed by HRP and AP convert chromogenic substrates such 3,3',5,5'tetramethylbenzidine (TMB) and 2,2'-Azinobis [3-ethylbenzothiazoline-6-sulfonic acid]-diammonium salt (ABTS) into colored products due to their high catalytic activity and specificity.^{103–106} While each detection enzyme can amplify the signal by transforming $10^3 - 10^4$ molecules of the substrate into product per minute²⁴ the binding of HRP or AP directly to detection antibody results in only a 1:1 stoichiometric ratio between enzyme and detection antibody.¹⁰⁷ Increasing this ratio for enhancement of signal produced per detection antibody bound has been achieved by exploiting the relationship between the molecules streptavidin and biotin, which act as a bridging system to enable multiple signal producing enzymes to bind to a single detection antibody. Streptavidin is a bacterial protein tetramer that contains four high-affinity binding sites for biotin, a B vitamin. Detection antibody that has been biotinylated at multiple sites will tightly bind multiple enzyme-streptavidin conjugates, thus amplifying the signal produced from a single bound detection antibody. While this classically used system is successful in measuring analytes as low as 0.01 ng mL^{-1, 9,10} there is a clinical need to reliably detect disease associated biomarkers that typically exist below the limits of this method. Modifications of this approach have generally involved increasing detection enzymes per

detection antibody bound to further amplify the signal. This is achieved with the Tyramide amplification system which processes biotinylated Tyramide in the proximity of detection antibody to adhere more HRP enzymes and thus boost the amount of substrate processed per detection antibody.^{108,109} In general, Tyramide amplification and other related approaches do provide appreciable boosts in sensitivity but fall short of reliable detection at sub-nanomolar concentrations.

Nanoparticles, particularly gold (GNP) and magnetic nanoparticles (MNP), have been incorporated into ELISA platforms in several different roles, including both as substrate replacements and as multivalent shuttles for detection antibody and enzymes to enhance the signal. GNPs have become the preferred nanoparticle due to their simpler synthesis and highly modifiable surface chemistry.¹¹⁰ Integration of nanoparticles in various roles within ELISA has been shown to increase sensitivity and has been a source of great interest in the pursuit of reliable assays for the detection of trace analytes that lie below the limits of detection for traditional ELISA. Plasmonic ELISA takes advantage of the propensity for the surface electrons of noble metal nanoparticles such as gold and silver to oscillate and produce light absorption and reflection properties in the visible and near-infrared range when excited by light^{111–114} as shown in Figure 3. This characteristic oscillation, known as localized surface plasmon resonance (LSPR), and its visible color output can vary by nanoparticle size, shape, and interparticle proximity.¹¹⁵⁻¹¹⁷

Nanoparticles have been integrated into ELISA as detection systems by using enzyme-driven reactions to alter the proximity, physical attributes, or environment of nanoparticles in solution. This drives a change in LSPR

and thus a visible color change that is a function of the analyte concentration.¹¹⁴ For example, acetylcholinesterase enzyme (AChE) has been utilized to alter the LSPR of GNPs to induce their aggregation in proportion to the bound analyte.^{118,119} During detection steps, the substrate for AChE, acetylthiocholine, is added to the sample plate well along with citrate-capped GNPs, and the subsequent hydrolysis of acetylthiocholine by AChE produces cationic thiocholine. This positively charged thiocholine covalently binds to the negatively charged GNPs, causing a shift in charge distribution and subsequent aggregation and color change from red to blue reflecting analyte concentration. Aggregation-based formats can be highly sensitive, up to a thousand-fold more sensitive than traditional ELISA.¹¹⁸ However, a major limitation is the sensitivity of this system to spontaneous aggregation due to environmental factors, which has limited its real-world application.¹²⁰ Additional detection approaches that alter LSPR on the nanoparticle surface have included altering the formation and growth dynamics of gold nanoparticles from gold salt, which is driven by the concentration of reducing agent, which is consumed by the bound detection enzyme. This strategy was used to develop an ELISA against prostatespecific antigen (PSA) and human immunodeficiency virus-1 (HIV-1) capsid protein with the detection enzyme catalase, which consumes the reducing agent H₂O₂. After the formation of the immunosandwich, a solution of H_2O_2 and gold salt are added and in the absence of analyte the H₂O₂ concentration remains high, rapidly reducing the gold salt into non-aggregated GNPs, resulting in red color. Growth is subsequently slowed when analyte and thus detection antibody-bound catalase are present,



Figure 3 Schematic representation of plasmonic ELISA.

resulting in reduced H_2O_2 , slow growth of aggregated nanoparticles, and a blue color.¹¹ This ultrasensitive technique provided a limit of detection as low as 1 attogram mL⁻¹ of PSA or HIV-1 capsid, which could be observed with the naked eye. Plasmonic ELISA is a highly sensitive detection and signal amplification approach and although it is not without its drawbacks, namely environmentally induced aggregation producing non-specific signal, the ability to observe the presence of antigen with the naked eye makes it a promising platform for point-of-care (POC) devices.

Utilizing GNPs as scaffolds to conjugate multiple signal-generating enzymes per detection antibody takes advantage of the high surface area to volume ratio and tunable surface chemistries inherent to nanoparticles. For example, Duan et al directly adsorbed HRP-labeled detection antibody onto GNPs to quantify the amount of nucleocapsid protein (NP) of the novel bunyavirus for diagnostic purposes. The subsequent increase in sensitivity (limit of detection 0.9 pg mL⁻¹) was significantly higher than traditional ELISA and could even detect protein NP more sensitively than quantitative real time-PCR (qRT-PCR) could with NP mRNA.¹³ This concept has been taken even further with the coupling of polymers and dendrimers to the surface of the GNPs to serve as additional scaffolding to add yet more binding sites for enzymes and antibodies. Modifying GNPs with polyamidoamine (PAMAM) dendrimer has been used to conjugate up to 20 HRP enzymes on a single particle, producing significant signal amplification that results in double the sensitivity over traditional ELISA in detecting human chorionic gonadotropin.14

Significant signal amplification over traditional ELISA has also been attained by utilizing MNPs as vessels for the delivery of antibodies and HRP enzymes. Wu et al used this approach to magnetically enrich *Escherichia coli* from river water samples by employing MNPs coated in poly-L-lysine (PLL) brushes, which have copious amine groups for the binding of antibody and HRP.¹²¹ The magnetic purification coupled with the signal enhancement of multi-HRP enzyme binding to each MNP resulted in a limit of detection 40-fold lower than traditional ELISA.

Nanoparticles have also been investigated as replacements for enzymes such as HRP and AP in ELISA. Nanoparticles with enzyme-like properties, so-called nanozymes,¹²² were first described in 2007 with the reporting of a Fe₃MnO₄ NP that has peroxidase-like activity and since then nanozymes that mimic the activity of oxidase, catalase, and superoxide dismutase have also been discovered.^{122–128} Nanoparticles often have a lower cost and higher stability than traditional ELISA signal amplification enzymes and have been shown to be more sensitive. Ye et al demonstrated that an enzyme-free ELISA that instead utilized gold vesicles encapsulated with Palladium–Iridium bimetallic nanoparticles as peroxidase mimics was able to achieve a limit of detection in the femtogram mL⁻¹ range, 10³ lower than traditional ELISA.¹²⁹

DNA-based amplification strategies have had success in producing ultrasensitive ELISAs that can detect the antigen in the sub-femtomolar range. Immuno-PCR hybrid assays link the specificity of ELISA immunoassays with the amplification capability of PCR by employing a DNA-tag to mark the ELISA detection antibody which is subsequently amplified and detected through either gel electrophoresis or more commonly qRT-PCR analysis.¹² The DNA-tag is typically linked to the detection antibody directly or through a linker such as the commonly utilized biotin-streptavidin or several different covalent conjugates. Linking of DNA-tag and antibody, regardless of method, can be a complicated procedure and a thorough evaluation must be carried out to ensure that antibody affinity has not been compromised.¹³⁰ The signal can be further enhanced by biotinylating the DNA-tag at both ends, resulting in the formation of supramolecular complexes linked through streptavidin, with potentially hundreds of DNA-tags per detection antibody to further increases sensitivity.131

While immuno-PCR is 10⁵-fold more sensitive than standard enzyme-conjugated amplification, the background signal is increased due to the ability of PCR to amplify even a single DNA molecule that has adhered due to a nonspecific interaction. Reducing the background signal of the immuno-PCR platform is a crucial optimization point for the assay to reach its full sensitivity potential and several approaches have been developed, including the proximity ligation assay (PLA) to mitigate this issue.¹³²⁻¹³⁵ PLA employs two antibodies reactive to two different epitopes on the same antigen target, which have been conjugated with different DNA-tags. When the antigen target is present in a patient sample, the two antibodies along with their DNA-tags, are brought into proximity of one another enabling their ligation by added DNA ligase. This newly fused DNA-tag then acts as a template for rolling circle amplification, producing a highly sensitive assay with exceptionally low background due to the low odds of nonspecific interactions bringing the two different DNA-tags close enough for ligation.¹³⁶

PCR amplification of the DNA-tag requires temperature fluctuations that, while essential for successful transcription, can be detrimental to the stability of protein components in the immunoassay, as evidenced by the observation that when all ELISA and PCR steps are carried out in the same physical space, there is a 1000-fold decrease in sensitivity.^{137–139} One solution is to physically segregate the initial immunoassay and ensuing PCR steps by using a DNA-tag that can be cleaved from bound detection antibody at the end of the ELISA procedure, enabling the isolation of the relevant DNA-tag and performing subsequent PCR steps in a separate tube, significantly improving sensitivity.139 Isothermal amplification systems have also been developed to address this reliability issue and enable the transcription of DNA template without temperature cycling. Fluorescent amplification catalyzed by T7 polymerase technique (FACTT) isothermally generates RNA from a DNA-tag consisting of a tetrameric streptavidin molecule with 3 of 4 biotin-binding sites occupied by the DNA template, leaving the fourth to adhere to biotinylated detection antibody as shown in Figure 4.15 RNA transcribed by T7 polymerase is then visualized by the fluorescent RNA intercalating dve RiboGreen and quantified by a fluorometer, which improves sensitivity 10⁵ over traditional ELISA and has enabled detection of protein targets as low as 0.08fM.¹⁵

Like FACTT, the recently developed immunoassay coupled with isothermal exponential amplification reaction (IMEXPAR) employs a detection antibody labeled with a DNA primer using a biotin-streptavidin linker.¹⁴⁰ After the immunosandwich is formed, a DNA amplification template, DNA polymerase and nicking enzyme are added to mediate an exponential cycle of replication that is proportional to bound detection antibody and can be quantified with the use of the fluorescent nucleic acid dye SYBER Green I and qRT-PCR to a lower limit of detection of 1.63pM.

Enhancing immuno-PCR with GNPs or MNPs has also been shown to be a highly sensitive method of signal amplification. The employment of these particles in a bifunctional manner, in which they are coated with both detection antibody and DNA-tag, is an effective way to avoid the negative effects on antibody affinity of direct detection antibody/ DNA-tag conjugation.¹³⁰ Furthermore, GNPs have even been manipulated into a "polyclonal" status, wherein the nanoparticle has been coated with detection antibodies that react to different epitopes on the same antigenic target, enhancing their ability to bind the antigen target.¹⁴¹ Nam et al developed a biobarcode system that incorporates both GNPs and MNPs, resulting in a highly specific and sensitive immuno-PCR assay that has a reported limit of detection as low as 0.1 fg mL⁻¹.^{142,143} MNPs coated with capture antibody are used to magnetically isolate the immunosandwich that is subsequently formed with detection antibody and DNA-tag coated GNPs, after which the DNA-tag is dissociated from their GNP scaffold and analyzed by qPCR. Table 2 summarizes different methods for detection of analytes and signal amplification methods used.

Improving the amplification of ELISA signal has been a decades long pursuit and has resulted in both modifications



Figure 4 Schematic representation of FACTT. Reproduced from Zhang H, Cheng X, Richter M, Greene MI. A sensitive and high-throughput assay to detect low-abundance proteins in serum. Nat Med. 2006;12(4):473–477.¹⁵

S.No.	Target	Detection Method Employed	Amplification Method Used	Detection Limit	Reference
1.	Prostate Specific Antigen (PSA) and HIV-1 Capsid antigen	Plasmonic ELISA	Gold nanoparticles (Au NP)	I × 10 ⁻¹⁸ g/mL	[1]
2	Bovine Serum Albumin (BSA)	Immuno-PCR	Streptavidin-Protein A chimera attached to biotinylated DNA was amplified by PCR and detected by ethidium bromide staining	260bp fragment observed with 580 antigen molecules (9.6 × [10. ⁻²²] mol)	[12]
3.	Nucleocapsid protein of SFTSV	Gold nanoparticle-based ELISA	Au NP labeled with HRP-labeled monoclonal antibody	0.9pg/mL	[13]
4.	hCG	Gold nanoparticle-based ELISA	Assembly of Au NP induced by functional polyamidoamine dendrimers	0.03IU/L	[14]
5.	HER2	FACTT (Fluorescent Amplification catalyzed by T7 polymerase Technique)	T7 RNA polymerase amplifies the DNA template in the streptavidin-biotin- dsDNA template module bound to Ag- Ab complex	0.08fM	[15]
6.	Gliadin (in Celiac disease)	Indirect ELISA using Au NP as probes	Au NP labeled with HRP-labeled antibody	180pg/mL	[110]
7.	E. coli 0157:H7	Magnetic ELISA	Poly-L-Lysine mediated brushes on Magnetic NP (MNP) binding to Ab-HRP	8cfu/mL	[121]
8.	Prostate Surface Antigen (PSA)	Enzyme free ELISA	Enzyme free signal amplification based on gold vesicles encapsulated with Pd-Ir NP	3 l fg/mL	[129]
9.	Prostate Surface Antigen (PSA)	Loop mediated isothermal amplification (LAMP) in combination with Solid Phase- Proximity Ligation Assay (SP-PLA)	LAMP	0.0001pM	[135]
9.	Prostate Surface Antigen (PSA)	Immuno PCR in combination with Solid Phase-Proximity Ligation Assay (SP-PLA)	Real time PCR	0.1pM	[135]
10.	Shiga Toxin producing E. coli	Immuno-PCR	Immuno capture of Shiga toxin 2 and amplification of a DNA marker using real time PCR	0.1pg/mL	[137]
11.	MUCI (tumor protein Mucin I)	Sandwich immunoassay coupled to EXPAR (isothermal exponential amplification reaction)	EXPAR signal amplification and real- time quantitative PCR (rt-qPCR)	I.63pM	[140]
12.	Respiratory Syncytial virus (RSV)	Nanoparticle amplified-Immuno PCR (NPA-IPCR)	Signal amplification using multivalent Au NP followed by real time PCR	4.1 PFU/mL	[143]

to traditional ELISA as well as entirely new assays that combine previously uncoupled techniques to detect biomarkers in trace amounts. Undoubtedly, some of the lowest detection limits have been recorded using nucleic acid-based amplification techniques. However, these immuno-PCR platforms are not without drawbacks, including the length of the procedure, which can take up to two days.¹⁴⁴ Continued improvements in ready-made components of the immuno-PCR system, as well as the incorporation of nanoparticles, are now needed to reduce the total assay time from start to finish.

Discussion/Conclusion

ELISA has been extensively used to assay for proteins of interest in both clinical and research applications and improvements to the traditional ELISA platform will be necessary to maintain clinical relevance. The sensitivity of immunoassays is primarily dependent on the characteristics of the assay antibodies and evolving techniques to generate epitope-targeted antibodies and their constituent fragments provide the pathway to develop highly sensitive assays free from any interfering components. Furthermore, as more disease-related biomarkers are discovered to be clinically relevant in the sub-nanomolar range, ultrasensitive amplification techniques such as those reviewed here will need to be employed.

Typical antibody validation protocols seek to establish candidate antibodies as specific, selective, and reproducible in the intended application and the continued growth of mAb therapy provides a strong justification to incorporate epitope mapping in validation protocols for diagnostic antibodies. Characterization of ELISA antibodies has traditionally been relatively opaque, with commercial diagnostic antibodies rarely having defined or disclosed epitopes. Academic literature is rife with reports on the inadequacy of commercial antibody validation, 145-147 and this is particularly troubling when immunoassays are designed to evaluate antigens that may already have a therapeutic antibody bound, as is often the case during HER2 ECD quantification by ELISA in HER2-positive breast cancer. Since the cloning of the rat HER2 gene neu¹⁴⁸ and subsequent generation of a mouse mAb reactive to its ECD,¹⁴⁹ highly impactful targeted antibody therapies with defined epitopes such as trastuzumab in subdomain IV and pertuzumab in subdomain II have been generated, characterized, and extensively administered in HER2-positive breast cancer patients.⁵² The majority of diagnostic HER2 antibodies have not been similarly classified, however. Recent reports are recognizing the potential peril of targeted HER2 mAb therapy interference in ELISA,³⁹ while justifiably advocating for the mapping of diagnostic antibody epitopes using a characterization pipeline that includes testing for therapeutic mAb interference. These approaches are essential for the development of an unbiased, reliable evaluation of therapeutic efficacy to inform clinical treatment decisions during targeted mAb therapy.

As disease-related biomarkers continue to be discovered at widely varying concentrations within biofluid, the need for signal amplification techniques will become ever more important to meet the challenge of ultrasensitive detection. Biomarkers such as programmed death-ligand 1 (PD-L1), a transmembrane ligand that suppresses the adaptive immune system through binding to its receptor PD-1,¹⁵⁰ have been implicated in nonsmall cell lung carcinoma^{151,152} and breast cancer¹⁵³ at sub-ng mL-1 concentrations, below the reliable detection limits of a traditional colorimetric ELISA. Similarly, earlier detection of biomarkers such as cardiac troponin I, which are only detectable in blood during pathological conditions such as acute myocardial infarction,^{154–156} are enabled by more sensitive techniques that have lower limits of detection, providing opportunities for earlier intervention and potentially improved clinical outcomes.

Ideally, amplification techniques should be facile and transferable into POC devices such as lateral-flow assays (LFA) or microfluidics platforms. Indeed, commercial LFA devices are already employing gold plasmonics and magnetic nanoparticles as stable and sensitive methods of signal amplification.¹⁵⁷ These devices have the potential to be particularly transformative in underserved regions of the world, where the World Health Organization estimates only 35% of low-income countries have accessible pathology services for diagnosis.¹⁵⁸ Targeted treatment for breast cancer is dependent upon biomarker analysis that is traditionally performed by pathologists on tumor biopsies. For example, gualification for targeted mAb therapies trastuzumab and pertuzumab relies upon demonstration of HER2 overexpression by either IHC or fluorescent in situ hybridization (FISH), analyses often unavailable in resource constrained locales. While tissue HER2 analysis is the gold standard for diagnosing HER2-positive breast cancer, soluble HER2 ECD quantification could effectively bring underserved patients off the sidelines by diagnosing those that have an elevated serum HER2 ECD level at the time of presentation and enabling access to targeted therapies that have proven outcome benefits.

Much progress has been made in enhancing the sensitivity, and thus utility, of the traditional ELISA format. As more proof-of-concept improvements migrate from the research laboratory to the clinical diagnostic setting, reliable sub-nanomolar detection capabilities will enable the rapid quantification of trace biomarkers for earlier diagnosis and effective treatment management.

Author Contributions

All authors contributed to drafting or revising the article, gave final approval of the version to be published, agreed to the submitted journal, and agreed to be accountable for all aspects of the work.

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Disclosure

Dr Hongtao Zhang owns stocks in Martell Diagnostic Laboratories, outside the submitted work; In addition, Dr Hongtao Zhang has patents Greene, MI, Eberwine, J, Kacharmina, JE, and Zhang, H-T. Methods, systems and kits for immuno-detection of epitopes expressed on molecules. US 6,743,592); Greene, M. I., Zhang, H. Methods of detecting molecules expressing selected epitopes via fluorescent dyes. US 7,045,286; Greene, MI, Eberwine, J, Kacharmina, JE, and Zhang, H-T. Methods for immunodetection of epitopes. US 7,341,831 issued May 16, 2006; Greene, MI, Eberwine, J, Kacharmina, JE, and Zhang, H-T. Method, system and kits for immuno-detection of epitopes expressed on molecules. US 7,361,464; M. I., Zhang, H. Methods of detecting molecules expressing selected epitopes via fluorescent dyes. US 7,524,628 issued related to FACTT mentioned in the review. Dr Franklin Pass is an employee of Martell Diagnostic Laboratories. Dr Mark I Greene reports grants from Martell, during the conduct of the study; grants from the NIH and BCRF, outside the submitted work; In addition, Dr Mark I Greene has a patent on erbB2 therapy owned by the University of Pennsylvania licensed to Roche-Genentech. The authors report no other conflicts of interest in this work.

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