#### **REVIEW**



# **Point‑of‑care electrochemical testing of biomarkers involved in infammatory and infammatory‑associated medical conditions**

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# **Abstract**

Recent years have shown that the diagnosis and monitoring of biomarkers involved in inflammatory-associated medical conditions such as cancer, neurological disorders, viral infections, or daily physical activities offer real benefits in increasing the quality of medical care and patient life quality. In this context, the use of integrated and portable platforms as point-of-care testing devices for biomedical analysis to enable early disease diagnosis and monitoring, which can be successfully used even at the patient's bed, is an emergency nowadays. The development of low-cost, miniaturized, and portable, user-friendly devices that provide an answer in a timely manner, such as electrochemical sensors, is relevant for the elaboration of point-of-care testing devices. This review focuses on the recent progress in bioanalysis of both specific biomarkers and inflammatory-associated biomarkers present in several diseases like neoplasia, severe neurological disorders, viral infections, and usual physical activity and provides an overview of the state of the art over the most recent electrochemical (bio)sensors for the detection of inflammation-related biomarkers. Future perspectives of point-of-care testing to improve healthcare management are also discussed.

**Keywords** Point-of-care testing · Biomarkers · Infammation · Electroanalytical methods · Wearable sensors

# **Introduction**

Infammation is a defense process of the body triggered by tissue damage. The etiology of infammation is diverse and can be represented by physical factors (trauma, extreme temperatures), chemical factors (endogenous or exogenous acids and bases), or infectious factors (bacteria, viruses, fungi, parasites). The infammatory process can be acute, accompanied by heat, redness, swelling, pain, and loss of function, or chronic when the processes of fbrosis and repair also occur.

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In acute infammation, the following molecules and systems are involved:

- 1. *Vasoactive amin*es: histamine, serotonin, involved in vasodilation and increase vascular permeability;
- 2. *Kinine*: bradykinin, involved in vascular changes, and pain;
- 3. *Complement system*: involved in immune defense and infammation;
- 4. *Coagulation: fbrinolysis system*: fbrin network blocks foreign particles;
- 5. *Mediators formed in the arachidonic acid cascades*: prostaglandins, prostacyclins, leukotrienes, and thromboxane, involved in pain, fever or infammation, and vascular processes;
- 6. *PAF-platelet activation factor*: involved in activation of thrombocytes and platelet aggregation;
- 7. *Reactive oxygen species*: superoxide anion,  $H_2O_2$ , hydroxyl radical, singlet oxygen;
- 8. *Reactive nitrogen species*: nitric oxide;
- 9. *Cytokines*: interleukins: IL-1, IL-6, IL-8; tumor necrosis factors: TNF-α, TNF-β; interferons: IFN-α, β, γ; colony-

stimulating factors G, M, GM: involved in pain and vascular and chemotactic processes [\[1](#page-24-0), [2](#page-24-1)].

In the case of chronic infammation, the aforementioned signs are not present, but the fnal consequences can be fbrosis, tissue destruction, and eventually necrosis. The cells and molecules involved in chronic infammation are monocytes, macrophages, lymphocytes, and cytokines. Cytokines are protein molecules that intervene in intercellular signaling and participate in cell growth and differentiation, apoptosis, and appearance and maintenance of infammation, respectively in angiogenesis. Regarding infammation, IL-6 and TNF-alpha are the most wellknown cytokines with a pro-infammatory role. IL-6 supports infammation through the secretion of cytokines and monocytes as well as through the activation of signaling pathways involved in tissue destruction. TNF-alpha supports and promotes infammation by stimulating the secretion of some proteins and immune cells and by stimulating coagulation (which can lead to occlusion or disseminated intravascular coagulation in case of severe infammation) [[3\]](#page-24-2). Because prolonged inflammatory processes have serious consequences on the patient's health and can lead to the appearance or aggravation of other pathologies, its early diagnosis is of interest. All these molecules can be used as fuid biomarkers for the efective diagnosis of infammation [[1,](#page-24-0) [2](#page-24-1)].

A biomarker is a molecule (i.e., a protein) or a biological process (alterations of nucleic acids) that can be identified and quantified to help diagnose and monitor a physiological/pathological process. Depending on their clinical role, biomarkers can be classified into diagnostic, monitoring, response, predictive, risk, and safety biomarkers [[4](#page-24-3)]. Biomarkers are true indicators of pharmacological reactions to a therapeutic strategy, thus being important to find effective strategies for early-stage diagnosis and for monitoring the treatment of different medical conditions [[5\]](#page-24-4). Also, some diseases, such as cancer or neurodegenerative diseases, severely affect the general condition of the patient and are potentially fatal; thus, it is important to administer a specific treatment as early as possible  $[6]$  $[6]$  $[6]$ .

Several medical conditions are accompanied by a chronic infammatory process that can trigger the activation of other processes such as tissue destruction or degeneration, and the release of metabolites with local or systemic toxicity, respectively. Besides the specifc markers of infammation, the products determined by infammatory processes can also be considered biomarkers. This is mainly due to the fact that infammatory biomarkers do not provide information on the location of the infammation or the afected organ, respectively, of the actual diagnosis. Thus, it is necessary to correlate the markers of infammation with those characteristic of that condition [[7](#page-24-6)].

According to the International Standard ISO22870, *Pointof-care testing (POCT): Requirements for quality and competence*, a POCT is a test "performed near or at the site of a patient with the result leading to a possible change in the care of the patient" [\[8](#page-24-7)]. A POCT can be performed in different locations such as at home, pharmacies, ambulances, nursing homes, physician's offices, remote medical centers, hospitals, sports clinics, or workplaces. POC are analytical devices such as test kits or glucometers that are provided for in situ testing of the patient/user rather than sample collection and analysis in a traditional environment such as a clinical laboratory.

Biosensors are analytical devices composed of a transducer, bioreceptor, and digital output detector that can convert a biological or biochemical event into a measurable signal via the transducer. Depending on the type of the detection method, the signal is generated directly at the transducer surface (impedance or surface plasmon resonance) or indirectly by means of signaling molecules, such as fluorophores, enzymes, and electrochemically active molecules [[9](#page-24-8)]. Based on the signal from the biorecognition reaction that undergoes transduction, the electrochemical biosensor market is segmented into potentiometric, impedimetric, conductometric, voltammetric, and others [[10](#page-24-9)].

In this review, infammation markers involved in diseases characterized by infammatory processes such as cancer, neurological and neurodegenerative diseases, and SARS-CoV-2 infection (COVID-19) are presented. Moreover, examples of markers associated with infammation produced by sports activities are also discussed. An overview of the electrochemical detection methods for these types of biomarkers and the ones suitable for POCT was discussed in the following sections. Figure [1](#page-2-0) outlines the main directions approached in this review, highlighting the specifc characteristics of POCT.

# **Point‑of‑care electrochemical testing**

The development of new methods for POCT using streamlined and ideally reagent-free assays is a timely, critical, but challenging issue. Although reference methods achieve clinically relevant results in terms of accuracy, sensitivity, and selectivity, they oftentimes are lengthy, expensive, and require laboratory settings [[11](#page-24-10)].

Recently, electrochemical methods have received increased attention due to advantages such as fast response, low use of reagents, and high selectivity and sensitivity, making them suitable for custom-made sensors [[12\]](#page-24-11). A fast search on the ScienceDirect platform by using "point-ofcare detection" as a keyword has generated 365,181 results,



<span id="page-2-0"></span>**Fig. 1** General illustration of the use of POCT devices for diagnosis, monitoring, or prognosis purposes in infammatory-related diseases for improved healthcare management (original fgure created with Biorender.com)

including articles, reviews, and book chapters. By adding "electrochemical" as a supplementary keyword, 21,464 results were generated, which is roughly 6%, highlighting the increased interest in this topic.

Several features of electrochemical (bio)sensors must be considered for efficient POCT device development (Fig. [2\)](#page-3-0): new electrochemical substrates and simple detection methods, nanomaterials for increased sensitivity and reduced biofouling of targets, bioreceptors for increased specifcity and multiplexed analyses, easy powering and low-cost electrochemical analyzers, digital outputs, and the combination with other technologies [[13–](#page-24-12)[16\]](#page-24-13).

Two diferent approaches in the feld of POCT emerge from the literature with the involvement of electrochemical biosensors: the miniaturization of components, to ensure portability, and the integration of bioelements to ensure selectivity for the target. In addition, it is very important that the system be tested in clinical-related settings. There are regulations imposed by the World Health Organization (WHO) in this regard, according to which the systems can be classifed as POCT only if they meet all the conditions included in the ASSURED guidelines: "Affordable," "Sensitive," "Specific," "User-friendly," "Robust and rapid," "Equipment-free," "Deliverable" to all people needing the test, including low-resource communities [\[17\]](#page-24-14).

Some of the molecules determined by using POCT devices are represented by functional nucleic acids (DNA, RNA, aptamers), proteins (cytokines, enzymes, and antibodies), hormones, toxins, infectious particles (viruses and bacteria), and cells. The main purpose of the elaboration of biosensors is to specifcally detect these molecules at their very low levels found in complex biological matrices. An important category of POCT devices applied in the biomedical feld is based on antibodies and aptamers as modifers for highly selective transducers [[18,](#page-24-15) [19\]](#page-24-16). The analytical performances of these devices can be increased by using the new fndings and developments in nanomaterials, in chemical and biological domains. Furthermore, the outstanding progress



<span id="page-3-0"></span>**Fig. 2** Schematic representation of POCT electrochemical devices (original fgure created with Biorender.com)

accomplished in miniaturization, electronics, artifcial intelligence (AI), and the Internet of Things (IoT) allowed the development of wearable and portable (bio)sensors suitable for POCT device fabrication [\[11](#page-24-10), [17](#page-24-14)].

# **Biomarkers involved in infammatory diseases**

### **Cancer biomarkers**

#### **Medical aspects related to cancer biomarkers**

Cancer biomarkers can be classifed into two major categories: molecular and processes (apoptosis, angiogenesis, and proliferation) biomarkers. The molecular biomarkers comprise *nucleic acids* (include modifcation of the entire genome and of genes primary sequence, such as rearrangements, amplifcations, deletions, fragmentation of DNA/ RNA), *proteins* (Bence-Jones proteins, carcinoembryonic antigen (CEA), cancer antigen-125 (CA-125), prostate-specifc antigen (PSA), ApoA1, human epididymal secretory protein 4 (HER4), β-2 microglobulin, programmed death ligand1 (PD-L1), hypoxia-inducible factor1alpha (HIF-1 $\alpha$ ), tumor-associated and specific antigens), *metabolites* (fumarate, succinate, D-2-hydroxyglutarate (D-2HG);  $H_2O_2$ ), *exosomes* (glypican-1 (GPC1), CD9, CD81, GTPases, miRNA), and *circulating tumor cells* (CTCs)[[14](#page-24-17), [20–](#page-24-18)[26](#page-24-19)].

Cancer biomarkers are secreted into tissues and biological fuids, allowing early detection by minimally invasive and non-invasive methods [[27,](#page-24-20) [28](#page-24-21)]. Here, we can include all analytical devices and methods that can be applied for the detection and/or monitoring of biomarkers from biological samples that can be collected in a minimally invasive manner (saliva, perspiration, tears), such as portable and wearable electrochemical detection devices. Any biomarkers in the category of infammatory factors such as tumor necrosis factor (TNF-α), interferon-gamma (IFN-γ), and interleukin-6 (IL-6) can be also considered biomarkers for cancer. Numerous studies in the literature are based on the detection of these compounds, for which an attempt is made to fnd a correlation between their levels in diferent biological fuids and the stage of the disease [[16\]](#page-24-13).

#### **POCT electrochemical sensors**

Due to the advantages of portable and wearable devices in biomedical applications, there is an increase in the number of studies focused on electrochemical platforms developed for the analysis of cancer biomarkers [\[14](#page-24-17), [21,](#page-24-22) [29\]](#page-24-23). Electrochemical portable devices for monitoring key biomarkers are the potential next frontier of wearable technologies for POCT diagnosis, including cancer. Furthermore, the extraordinary development in the feld of microfuidics allows the detection of traces of biomarkers in small volumes of biological samples because it involves a pre-concentration step before detection [\[15,](#page-24-24) [30](#page-24-25)]. Numerous approaches have been published regarding the detection of a large number of biomarkers expressed in cancers using miniaturized devices or sensors suitable for miniaturization [[15,](#page-24-24) [31](#page-24-26), [32](#page-25-0)]. The aspect related to miniaturization is decisive when the development of POCT or point-of-use (POU) devices for decentralized detection, in the feld, at the patient's bedside is aimed. Thus, easy to manufacture, and low-cost detection of cancer biomarkers is crucial for fghting early-stage disease and increasing the patient's chances of healing and survival. Some examples of such detection devices and their analytical parameters are presented in Table [1,](#page-5-0) the analytical methods being further discussed pointing out the most important components (nanomaterials and bioelements) and if real samples were analyzed.

The use of nanomaterials and composite materials for the functionalization of electrodes before the immobilization of biocomponents for the elaboration of electrochemical biosensors proved to be beneficial, especially in increasing the sensitivity for the detection of targets [\[35](#page-25-1), [36,](#page-25-2) [43](#page-25-3), [44,](#page-25-4) [46](#page-25-5)]. An example of such a material which is intensively used in the electrochemical analysis of biological compounds, including cancer biomarkers, is graphene. Graphene improves the analytical performance of platforms due to its excellent electrical and thermal conductivity, mechanical strength, large surface area, and the possibility of being further functionalized with various compounds or nanostructures [\[13,](#page-24-12) [40](#page-25-6), [69\]](#page-26-0). The development of portable (bio)sensors based on nanomaterials and composite materials is also related with the discovery of versatile materials, which in addition to special conductive properties, must be suitable for functionalization and structural handling. The methods by which graphene-based materials are transferred to the electrode are in continuous development, allowing to obtain high-purity, stable, and uniform flms via chemical vapor deposition (CVD) technique [[70](#page-26-1)] or laser-associated technique [[41,](#page-25-7) [71](#page-26-2)]. The functionalization of graphene with  $\cos_2$  and gold nanoparticles (AuNPs) was exploited for the modifcation of the surface of carbon-based SPEs, which allowed the detection and quantifcation of CA15-3 protein with good analytical performance, due to the excellent electrocatalytic activity of graphene-based immunosensor for the CA15-3 detection in serum employed the use of ternary Ag,  $TiO<sub>2</sub>$ , and reduced graphene oxide (rGO) to undergo signal amplifcation for the amperometric detection of  $H_2O_2$  promoted upon the antibody-antigen immunoreaction [[36](#page-25-2)]. Good analytical parameters were obtained for the detection of PSA using an ultra-sensitive electrochemical immunosensor based on AuNPs, p-aminothiophenol, and GO nanocomposite, and a self-assembled delaminated MXene-AuNP nanocomposite. By combining these nanomaterials, an increase in the amount of PSA antibodies captured at the electrode was obtained [\[49](#page-25-8)]. After the functionalization of the transducer with gold structure loaded on thionine functionalized GO layer, the immobilization of a high amount of antibodies and the acceleration of the electron transfer rate were both facilitated. This composite was further functionalized with PtCu bimetallic hybrid and loaded on 2D/2D rGO/graphitic carbon nitride, resulting in a complex material with excellent electrocatalytic properties, amplifed detection signal, and biocompatibility [[51\]](#page-25-9). Citrate-capped AgNP-modifed graphene QDs were synthesized, embedded in nano-ink, and applied as a starting platform for the development of an ultrasensitive sandwich-type portable device for the detection of PSA in raw plasma. Signal amplifcation was obtained using cysteamine functionalized with AuNPs [\[53](#page-25-10)]. An aerogel porous network nanocomposite was synthesized based on graphene functionalized with AuNPs and  $MoS<sub>2</sub>$ for the immobilization of a high amount of anti-PSA antibodies and the development of a label-free electrochemical immunosensor with good perspective for early diagnosis [[55\]](#page-25-11). A highly sensitive porous and highly biocompatible microgel nanocomposite based on AuNPs, polyacrylic acid, and N-isopropylacrylamide was used and allowed the covalent immobilization of the amino-modifed oligonucleotide capture probe complementary to the target. An optimized DPV procedure was applied to monitor the hybridization of miRNA-21 with the capture probe in serum [[72\]](#page-26-3).

the nanocomposite and increased amount of CA 15–3-specifc antibodies immobilized at the electrode [\[35\]](#page-25-1). Another

The category of carbon-based nanomaterial with special properties and which meets the necessary criteria in the development of electrochemical (bio)sensors also includes carbon nanotubes (CNTs). This material is suitable for functionalization with other materials, chemicals, or biocomponents, presenting many applications in the biomedical field. For example, the electrocatalytic activity of the magnetic framework  $Fe<sub>3</sub>O<sub>4</sub>@TMU-21$  multiwalled carbon nanotubes (MWCNT) was exploited for the construction of an electrochemical immunosensor for HER2 as a tumor biomarker in clinical monitoring. The immobilization of TMU-21 units at the surface of the  $Fe<sub>3</sub>O<sub>4</sub>$ -NPs was facilitated by methacrylic acid. The final composite material provided electrocatalytic activity

<span id="page-5-0"></span>





*CA 15–3* cancer antigen 15–3, *AuSPE* gold screen–printed electrode, *SWV* square wave voltammetry, *LSG* laser-scribed graphene, *AuNS* gold nanostructured, *MIP* molecular imprinted polymer, HER-2 human epidermal growth factor receptor 2, cTn-I cardiac troponin-I, cTn-T cardiac troponin-T, cTn-C cardiac troponin-C, Gluc glucose, DA dopamine, Chol cholesterol, Apr aptamer, *Au–Si-MNA* high-density gold-coated silicon microneedle arrays, *DPV* diferential pulse voltammetry, *BNC* bacterial nanocellulose, *CA-199* cancer antigen 199, *SPE* screen-printed electrodes, WVFY tryptophan-valine-phenylalanine-tyrosine, SP17 sperm protein 17, ITO indium tin oxide, APTMS alkoxysilane substituted polymer, MUCI mucin 1, IgG immunoglobulin G, BSA bovine serum albumin, *MWCNT-COOH* multi-walled carbon nanotubes, *GE* gold electrodes, *IL* interleukin, *CEA* carcinoembryonic antigen, *Her-2-ECD* extracellular domain of the human epidermal growth factor receptor 2, *DPASV* diferential pulse anodic stripping voltammetry, *QD* quantum dots, *SPCE* screen-printed carbon electrode, *GO* graphene oxide, *AuNPs* gold nanoparticles, protein, *GM2* ganglioside GM2 activator protein, *citrate AgNPs* citrate-capped silver nanoparticles, *GQDs* graphene quantum dots, *Cys-AuNPs* cysteamine with gold nanoparticles, *GIDE* gold interdigitated electrode, *HSA* human serum albumin, *hK2* human glandular kallikrein 2, *HE4* human epididymis protein 4, *MBs* magnetic beads, *SPCE Au NCs/MWCNT-NH2* gold nanocage graphene aerogels nanocomposite, *GCE* glassy carbon electrode, *AuPtAg ANCs* AuPtAg-alloyed nanocrystals, *CoS2-GR* cobalt sulfde/graphene nanocomposite, *GSPE* graphite screen–printed *141, let-7a* oligonucleotides, *PEI-Ag NPs* polyethyleneimine-silver nanoparticles, *HAS* human serum albumin, *HBS* hepatitis B surface antigen, *Fe3O4@TMU-21* magnetic framework, *PbS QDs* lead sulfde quantum dots, *MOFs* metal organic frame, *MB* methylene blue, *ATP* p-aminothiophenol, *d-Ti3C2TX MXene@AuNPs* delaminated MXene-gold nanoparticles, *CHI* chitosan,  $4\mu$ @Th/GO Au loaded on thionine-functionalized graphene oxide, PtCu@rGO/g-C3N4 PtCu bimetallic hybrid loaded on 2D/2D reduced graphene oxide/graphitic carbon nitride, Au@PDA@ BCN polydopamine-coated boron-doped carbon nitride, AuNPs-MG gold nanoparticles-magnetic graphene composite, CC/Co@C-CNTs Co-embedded N-doped carbon nanostructures, GQDs CA 15-3 cancer antigen 15-3, AuSPE gold screen-printed electrode, SWV square wave voltammetry, LSG laser-scribed graphene, AuNS gold nanostructured, MIP molecular imprinted polymer, HER-2 human epidermal growth factor receptor 2, cTn-1 cardiac troponin-1, cTn-T cardiac troponin-T, cTn-C cardiac troponin-C, Gluc glucose, DA dopamine, Chol cholesterol, Apt aptamer, 4u-Si-MNA high-density gold-coated silicon microneedle arrays, DPV differential pulse voltammetry, BNC bacterial nanocellulose, CA-199 cancer antigen 199, SPE screen-printed electrodes, WVFY tryptophan-valine-phenylalanine-tyrosine, SP17 sperm protein 17, ITO indium tin oxide, APTMS alkoxysilane substituted polymer, MUC1 mucin 1, IgG immunoglobulin G, BSA bovine serum albumin, MWCNT-COOH multi-walled carbon nanotubes, GE gold electrodes, IL interleukin, CEA carcinoembryonic antigen, Her-2-ECD extracellular domain of the human epidermal growth factor receptor 2, DPASV differential pulse anodic stripping voltammetry, QD quantum dots, SPCE screen-printed carbon electrode, GO graphene oxide, AuNPs gold nanoparticles, P3ABA/2D-MoSe2/GO poly(3-aminobenzylamine)/two-dimensional (2D) molybdenum selenide/graphene oxide, P3A prostate-specific antigen, UA uric acid, AA ascorbic acid, AFP alpha-fetoprotein, GM2 ganglioside GM2 activator protein, citrate AgMPs citrate-capped silver nanoparticles, GQDs graphene quantum dots, Cys-AuMPs cysteamine with gold nanoparticles, GIDE gold interdigitated electrode, HSA human serum albumin, hK2 human glandular kallikrein 2, HE4 human epididymis protein 4, MBs magnetic beads, SPCE Au NCs/MWCNT-NH2 gold nanocage graphene aerogels nanocomposite, GCE glassy carbon electrode, AuPAg ANCs AuPtAg-alloyed nanocrystals, CoS2-GR cobalt sulfide/graphene nanocomposite, GSPE graphite screen-printed electrodes, Au/MoS2/rGO gold nanoparticles/molybdenum disulfide/reduced graphene oxide nanocomposite, Ca chronoamperometry, CC cystain C, TB toluidine blue, miRNA-155, miRNA-141, let-7a oligonucleotides, PEI-Ag NPs polyethyleneimine-silver nanoparticles, HAS human serum albumin, HBS hepatitis B surface antigen, Fe3O4@TMU-21 magnetic framework, PbS QDs lead sulfide quantum dots, MOFs metal organic frame, MB methylene blue, ATP p-aminothiophenol, d-Ti3C2TX MXene@AuNPs delaminated MXene-gold nanoparticles, CHI chitosan, *BCN* polydopamine-coated boron-doped carbon nitride, *AuNPs-MG* gold nanoparticles–magnetic graphene composite, *CC/Co@C-CNTs* Co-embedded N-doped carbon nanostructures, *GQDs* graphene quantum dots, *AuPd-ANPs* AuPd alloy nanoparticles, CF carbon nanofiber, NCNTAs nitrogen-doped carbon nanotube arrays, PMMA-CD polymethyl methacrylate carbon dots, PPy graphene quantum dots, *AuPd-ANPs* AuPd alloy nanoparticles, *CF* carbon nanofber, *NCNTAs* nitrogen-doped carbon nanotube arrays, *PMMA-CD* polymethyl methacrylate carbon dots, *PPy P3ABA/2D-MoSe2/GO* poly(3-aminobenzylamine)/two-dimensional (2D) molybdenum selenide/graphene oxide, *PSA* prostate-specifc antigen, *UA* uric acid, *AA* ascorbic acid, *AFP* alpha-fetocoupled with an amidated multi-walled carbon nanotube, *IncRNAs* long non-coding RNAs, *HOTAIR*, H19, miRNA129 oligonucleotide sequences, AuNPs/MoS2-Gas Au nanoparticles/MoS2coupled with an amidated multi-walled carbon nanotube, *lncRNAs* long non-coding RNAs, *HOTAIR, H19, miRNA129* oligonucleotide sequences, *AuNPs/MoS2-Gas* Au nanoparticles/MoS2 electrodes, *Au/MoS2/rGO* gold nanoparticles/molybdenum disulfde/reduced graphene oxide nanocomposite, *Ca* chronoamperometry, *CC* cystatin C, *TB* toluidine blue, *miRNA-155, miRNA-Au@Th/GO* Au loaded on thionine-functionalized graphene oxide, *PtCu@rGO/g-C3N4* PtCu bimetallic hybrid loaded on 2D/2D reduced graphene oxide/graphitic carbon nitride, *Au@PDA@* polypyrrole, p-ABA p-aminobenzoic, pATP p-aminothiophenol, PPyr-NHS polypyrrole N-hydroxy succinimide polypyrrole, *p-ABA* p-aminobenzoic, *pATP* p-aminothiophenol, *PPyr-NHS* polypyrrole N-hydroxy succinimide toward the reduction of  $H_2O_2$  used for the indirect detection of the target in human serum [[39\]](#page-25-16). Hence, using Au nanocage coupled with an amidated MWCNT-functionalized SPEs enabled the efficient detection of lncRNAs in clinical settings [[73](#page-26-16)].

In addition to carbon-based materials, nanoparticles or nanostructures of noble metals are involved in the development of electrochemical biosensors. In the case of noble metal nanostructures, AuNPs and silver nanoparticles (AgNPs) represent the most common forms applied in this feld [\[50,](#page-25-21) [74](#page-26-17)]. In this case, the utility is related both to the improvement of the electrochemically active surface and implicitly of the sensitivity, and in facilitating the immobilization by strong and stable bonds of the structures with thiol group, especially in aptasensor development [[50\]](#page-25-21).

The use of antibody-type biological components in the fabrication of electrochemical immunosensors represented the starting point for the development of this feld ensuring promising perspectives for the biomedical domain. The presence of antibodies in the structure of electrochemical biosensors allowed the extraordinary improvement of the analytical performances in terms of selectivity for the target, this being mainly due to the specifc antigen–antibody interaction [[22,](#page-24-27) [33](#page-25-12), [35](#page-25-1), [39](#page-25-16), [46](#page-25-5), [62](#page-26-9)].

The most common strategy for immobilizing antibodies on the electrode surface is by amide, covalent bonding. For this purpose, carboxyl groups must be generated at the electrode to bind to amino groups in the antibody [[62,](#page-26-9) [63](#page-26-10)]. Thus, an innovative portable immunosensor based on a gold chip-type transducer functionalized with a self-assembled monolayer of mercaptosuccinic acid was designed for the detection of the CA 15–3. The covalent immobilization of the CA 15–3–specifc antibodies via EDC/NHS chemistry with the carboxyl groups provided by the polymer ensured the selectivity of the immunosensor for the target protein in serum [\[33](#page-25-12)]. In another study, antibody immobilization is performed by interaction with the functions present on the composite flm based on core/shell CdSe@ZnSQD, which also acts as an electroactive label, for the detection of HER2 in spiked serum [\[46](#page-25-5)]. A nano-biosensor for TNF- $\alpha$  was fabricated starting from a nanoflower-like structure of  $MoS<sub>2</sub>$ and applied for the evaluation of this biomarker in cancer patients' serum. The optimized biosensor showed excellent analytical performance, and good perspectives for application in decentralized analysis, being validated on cancer patient serum and by the standard ELISA method<sup>[[63\]](#page-26-10)</sup>.

Indirect amperometric detection via the reduction of  $H_2O_2$  in the presence of horseradish peroxidase (HRP) used as a label for the secondary antibody is the most common strategy for the detection of sandwich-type electrochemical immunosensors [[50,](#page-25-21) [75](#page-26-18)]. Conventional electrochemical immunosensors based on electroactive materials often sufer from leakage and signal instability. To overcome this issue, label-free detection was applied and, in this case, nanomaterials and redox samples play an important role in amplifying the signal [[47,](#page-25-19) [56,](#page-25-24) [63,](#page-26-10) [76\]](#page-26-19). Thus, an electrochemical immunosensor based on a fufy metal porous material, MOF-235, and methylene blue (MB) as an efective signal amplifcation strategy was designed and applied for the sensitive detection of PSA [[48\]](#page-25-20). A lead sulfde QDs-conjugated secondary HER2 antibody was used as a label. The indirect detection was obtained based on the SWV signal of the  $Pb^{2+}$  ions after their dissolution in HCl, enabling the detection of HER2 in serum with good recovery and selectivity [[42\]](#page-25-17).

Another fairly common strategy in recent decades for the design of electrochemical sensors is the use of biomimetic components, of which aptamers, microRNAs (miR-NAs), and molecularly imprinted polymers (MIPs) have proven very useful, especially in terms of lower cost and better stability compared to antibodies, without negatively infuencing the selectivity for the targets [\[23](#page-24-28), [41](#page-25-7), [54](#page-25-23), [66,](#page-26-13) [67,](#page-26-14) [77](#page-26-20)]. For example, a MIP-based electrochemical sensor was elaborated for CA 15–3 on AuSPEs. The imprinting process consisted of the preconcentration of the target protein on the electrode followed by the electropolymerization of the 2-aminophenol monomer. The extraction of protein after this step allowed to obtain of the MIP-type sensor, which was then tested by voltammetry in the presence of a redox probe [[37\]](#page-25-14). When poly(toluidine blue)-imprinted flm was produced for the same biomarker, a signal decrease of the redox probe in DPV was observed with increased protein concentration [[38\]](#page-25-15). The in situ electropolymerization was exploited for the generation of a MIP network consisting of poly(3,4-ethylenedioxythiophene) (PEDOT) and polypyrrole (PPy) at the surface of a carbon fabric anode containing Pt/Ru nanoparticles. This MIP-based sensor was designed as a simple, inexpensive, and efective tool for the detection of CEA in serum, being promising for further development of POCT [[45\]](#page-25-18). The combination of nanomaterials with MIP flms was also applied and has led to an improvement in parameters such as sensitivity, simplicity, porosity, and fexibility. Exploiting these advantages, a MIP sensor was developed using electrodes functionalized with graphene, nanostructured gold, and MIPs obtained after the electropolymerization. All these components embedded together in the flm deposited on the surface of the transducer allowed the selective and high-sensitivity detection of HER2in spiked undiluted serum [\[40](#page-25-6)].

Aptamers represent another category of biomimetic elements, diferent from MIPs, that are short, single-stranded DNAs or RNAs. Aptamers are versatile and bind targets with high selectivity and specificity since recognition and binding involve three-dimensional, shape-dependent, and hydrophobic interactions, base-stacking, and intercalation [[78](#page-26-21)]. The targets can range from small molecules, peptides, and carbohydrates to proteins or live cells, and bend into a variety of shapes due to their tendency to form helices and single-stranded loops. Being synthetic compounds, with superior stability and versatility to bioelements such as antibodies, aptamers have found numerous applications in the feld of electrochemical aptasensors [\[65,](#page-26-12) [79](#page-26-22)]. Thus, an aptamer with a high affinity for HER2 was immobilized onto the nanostructured electrode surface based on graphene and gold to enable the detection of HER2 in undiluted serum using an integrated handheld electrochemical system operated by a custom mobile application [[41](#page-25-7)]. Polymer flms (other than MIPs) deposited or generated on the electrode can also play an important role in the immobilization of the aptamer. Thus, p-aminobenzoic acid [[66](#page-26-13)], PPy [[67\]](#page-26-14), or a conjugated poly(pyrrole N-hydroxy succinimide) (PPyr-NHS) polymer [[68](#page-26-15)] are only some examples that have found their utility in the biomedical feld and were applied in the development of electrochemical aptasensors for IL-6 cytokine.

Detection of a single biomarker is usually not sufficient for accurate diagnosis. Thus, the selection of a panel of biomarkers that express themselves in the same condition would be extremely beneficial for the biomedical field. Such approaches have also been attempted with respect to the simultaneous detection of tumor biomarkers with portable POCT devices. For example, a dualmode electrochemical biosensor was developed for the simultaneous detection of two breast cancer biomarkers: CA15-3 and miRNA-21. The sensor composes of an SPE dual electrode array functionalized with a poly(3-aminobenzylamine)/two-dimensional/molybdenum selenide/ GO nanocomposite. The two working electrodes were functionalized individually with 2,3-diaminophenazine-AuNPs and toluidine blue (TB)–AuNPs. The redox probe-AuNPs were employed as signaling molecules and supports for immobilization of anti-CA15-3 antibodies and capture DNA-21 probes, also providing good selectivity, and sensitivity for the dual target detection, and great potential to be further developed for the simultaneous detection of other miRNAs and proteins for POCT diagnostic applications[\[34](#page-25-13)].

Some recent examples of electrochemical (bio)sensors for the detection of cancer biomarkers are presented comparatively in Table [1](#page-5-0).

#### **Neurological‑associated biomarkers**

# **Medical aspects related to neurological‑associated biomarkers**

Neural infammation is a defense mechanism of the organism which can represent a major symptom in many neurological disorders. Although acute infammation of the neural tissue may be beneficial, it can cause harm if the inflammatory reaction is prolonged or exacerbated. In some cases, tissue destruction can occur which can lead to irreversible degradation of the nervous tissue with the appearance of various forms of clinically manifest dysfunctions (i.e., epilepsy, Alzheimer's disease, Parkinson's disease, systemic lupus erythematosus, etc.) [[1,](#page-24-0) [80–](#page-26-23)[83\]](#page-26-24).

Among the biomarkers most frequently associated with neurological disorders are amyloid beta peptide (associated with Alzheimer's disease), the tau protein, and alphasynuclein (secreted in Parkinson's disease). Even if these biomarkers are specifc to certain neurological diseases, they can also be considered biomarkers associated with infammatory processes because they determine the occurrence of oxidative stress and other infammatory mechanisms (as seen in Table [2](#page-9-0)) [[84](#page-27-0)]. Biomarkers can be used for the rapid and efective diagnosis of neurological infammatory processes. As in the case of neoplastic diseases, biomarkers specifc for

<span id="page-9-0"></span>**Table 2** Examples of biomarkers involved in neurological disorders. Adapted after [[84](#page-27-0)]

Biomarker	Physiological implications	Pathological effect	Disorder
Amyloid beta	Lipid homeostasis, memory formation, regulation of synaptic activity, neural viability	Oxidative stress, synaptic dysfunc- tion, inflammation, apoptosis, neurite damage	Alzheimer's disease
Tau protein	Promotes microtubules polymerization, stability, and flexibility	Neuronal damage, pro-apoptotic	Alzheimer's disease, Parkinson's disease
Alpha-synuclein	Intervenes in the release of neuro- transmitters, modulates DNA repair processes	Synaptic and mitochondrial dysfunc- tions, oxidative stress, dysregulation of calcium signaling, microtubule impairment	Parkinson's disease
	Chromogramin A Vasoconstriction, regulation of glucose metabolism and storage, modulator of the neuroendocrine system	Oxidative stress, inflammation, neuronal Amyotrophic lateral sclerosis damage	
Micro-RNAs	Nervous system development and func- tion	Microglial activation, increase of proinflammatory cytokine release, and neuron loss	Autist spectrum disorders

infammation and non-specifc biomarkers which accompany infammation and concomitant processes (indirect determination) can be used [\[85](#page-27-1), [86](#page-27-2)]. A new category of biomarkers that have been recently introduced for diagnosis and monitoring of infammatory neurological disorders is represented by neuroflaments, the building blocks of proteins, and the basic structure of axons. The concentration of these proteins has been found to increase both in cerebrospinal fuid (CSF) and blood during neurological disorders and has been intensively studied. Neuroflaments can be used as biomarkers in neurological diseases such as stroke, traumatic brain injury, Parkinson's disease, multiple sclerosis, encephalitis, epilepsy, and peripheral neuropathies [\[87](#page-27-3), [88\]](#page-27-4).

The existence of validated markers for each action, but especially of fast, accessible methods of determination and allowing a decentralized use, at the patient's bed would be very benefcial for an early diagnosis of the disease. To date, numerous biomarkers have been reported correlated with neurological conditions. The most important of these are centralized in Table [2](#page-9-0), where the physiological implications, the pathological effect, and the type of disorder in which is expressed are also presented.

# **POCT electrochemical sensors**

Given that neurological diseases are often accompanied by infammatory processes whose consequences can be irreversible, early diagnosis of these diseases is of particular importance. There are many studies in this regard and the most recent will be discussed in this section of the review.

Based on the study of literature found on this topic, it is observed that the inclusion of nanocomposite materials has a crucial role in improving analytical parameters so the obtained sensors allow the detection of low concentrations of biomarkers when tested in complex matrix, as they are expressed in the neurological diseases for which rapid diagnosis is desired. The main advantage gained by the presence of nanomaterials in the sensor includes the existence of a greater number of active sites and functions available for immobilizing a larger number of bioelements and through stable interactions. This increases the sensitivity and stability of the (bio)sensors developed, catalyzes electrochemical reactions applied for the detection, enhances the electron transfer at the electrode, and assures the immobilization of diferent types of labels, which can highly improve the performance of the device [[13](#page-24-12), [89](#page-27-5)]. Thus, either graphene combined with metal nanoparticles [[89\]](#page-27-5), polymers combined with metallic foams [\[90\]](#page-27-6) or particles [[91](#page-27-7)], or composites based on noble metal particles or alloys [\[92\]](#page-27-8), or with MOFs [\[93\]](#page-27-9), were deposited on the electrode for increased sensitivity.

A small, portable magneto-immunosensor was developed for the amperometric detection of IL-6 by using a micro-fabricated electrode functionalized with rGO/AuNP nanocomposite to increase active surface useful for the immobilization of the antibody, fast charge transfer, and signal amplifcation [[30\]](#page-24-25). Streptavidin-inorganic hybrid nanofowers decorated with graphene were synthesized and used as substrates to modify GCEs and specifcally bind a diblock dual-aptamer allosteric hairpin, functionalized with AuNPs and a streptavidin-labeled aptamer for the detection of IFN- $γ$  [[94\]](#page-27-10).

The use of highly porous graphitic carbon deposited on polyimide tapes has proven useful in the development of a high-performance biosensor for IL-6 [[95\]](#page-27-11). A poly(curcumin-Ni foam)-based sensor was fabricated and successfully applied for the quantifcation of amyloid β oligomer (AβO), a circulating biomarker applied for early diagnosis of Alzheimer's disease [\[90\]](#page-27-6). For the same biomarker, another detection strategy was proposed using superhydrophobic carbon fber paper and AuPt-NPs to immobilize the aptamer strands specifc for AβO and indirectly minimize the nonspecifc adsorption effect due to its hydrophobic character [\[92](#page-27-8)]. After the functionalization of GCEs with cerium oxide nanofber electrospun, embedded in a Nafon layer, the composite flm generated at the electrode proved to be permeable in a selective way for the TNF- $\alpha$  biomarker, with fast electron transfer and excellent analytical performance in serum [[96](#page-27-12)].

Improving the analytical performance by using nanomaterials is not always adequate in the case of electrochemical sensors taking into account the principle on which their detection is based [[97\]](#page-27-13). Thus, in complex matrices such as real samples used to detect biomarkers involved in neurological disorders, the redox properties of the compounds simultaneously present in the sample can lead to mutual interference. To prevent this issue, the additional functionalization of nanostructured surfaces with bioelements and biomimetic elements is frequently applied strategies [\[75](#page-26-18), [98–](#page-27-14)[100](#page-27-15)]. An electrochemical immunosensor based on carbon SPEs functionalized with nanomaterials was designed for the accurate determination of unfolded p53 in plasma from patients diagnosed with mild cognitive impairment and Alzheimer's disease. The optimized detection method was validated by comparison with an ELISA assay, and the cheap, simple, and prone to miniaturization detection device has proved promising for POC applications [[101](#page-27-16)]. The presence of carboxyl groups provided by the 4-carboxymethylaniline substrates allowed the immobilization of the anti-TNF- $\alpha$  antibodies by an amide bond. A secondary antibody labeled with HRP was used for the detection, in which tetramethylbenzidine was also involved as a substrate for the electrochemical reaction. The optimized biosensor has been shown to be highly selective for the target when tested in the presence of other analytes secreted in the acute stage of infammation such as IL-10 and cortisol [[75](#page-26-18)]. The generation of amino groups on the transducer can also be used for the immobilization of

antibodies, since the latter have complementary carboxyl groups to achieve amide bonding [[9\]](#page-24-8). A polystyrene-polyamidoamine dendritic polymer nanofber–based platform was designed and provided a high density of amino groups useful for the covalent immobilization of a large number of capture antibodies with high selectivity for TNF- $\alpha$  in the presence of IL-10 and good perspective for clinical diagnosis [[102\]](#page-27-17).

Another strategy applied for the immobilization of biocomponents on the surface of the transducer is the esterifcation reaction in which the epoxy groups are involved. For example, the ultrasensitive detection of IL-6 has been achieved using a platform based on ITO substrate modifed with acetylene black and epoxy-substituted-poly(pyrrole) composite with good biocompatibility and conductivity. The high abundance of epoxy groups was useful for the esterifcation reaction applied for the reproducible and stable immobilization of the antibodies to promote the determination of IL-6 in samples from patients with neurological disorders [[103](#page-27-18)].

The use of aptamers as a biomimetic element for the selective detection of biomarkers involved in neuro-zone diseases has also been reported. Thus, a large number of AβO-specifc aptamers were immobilized on a composite layer based on CNF and AuPt alloy that allowed the elaboration of a high-performance electrochemical aptasensor for AβO [\[92\]](#page-27-8). The suitability of aptamers for functionalization allows the addition on their chain of some compounds or composites with a role in detection. For example, an innovative aptasensor was developed for IFN-γ based on targetinduced DNA strand and signal amplifcation with MB as a redox probe. It was observed a linear relationship between the signal of MB and IFN-γ content that could be successfully exploited for detection [\[91](#page-27-7)]. An aptamer-tethered gold nanofower was synthesized on the electrode, having a role in immobilizing both target capture and aptamer-tagged AuNPs/Cu-MOF conjugates. The use of MOFs as signal probes determined the amplifcation of the detected signal and the increase of the sensitivity for AβO electrochemical detection with a real perspective for the early diagnosis of Alzheimer's disease [[93\]](#page-27-9). The use of impedance as a detection method is another useful strategy that has found applications in the feld of aptasensors. A simple platform based on GO and AuNPs, embedded in a hydrogel layer, was used for the elaboration of an impedimetric label-free and selective detection of AβO in CSF or plasma [[89\]](#page-27-5). A label-free electrochemical aptasensor for TNF-α was fabricated based on a multifunctional DNA 4-way junction. The main advantage of this system was that it does not require additional labeling and signal amplifcation processes [[99\]](#page-27-19). An aptasensor for TNF- $\alpha$  was presented based on SPEs modified with gold hexacyanoferrate, AuNPs, and thiolated aptamers specifc for the target cytokine. The high sensitivity and wide detection range of this method allowed  $TNF-\alpha$  to be detected in serum even at very low concentrations as found in biological fluids [[104](#page-27-20)]. By combining an aptamer with silver nanoclusters and AuNPs immobilized on the amine-terminated electrode surface, a sensitive and specifc aptasensor for the detection of IFN-γ secreted by human cells was developed [[105\]](#page-27-21).

The use of platforms based on biomimetic elements, such as MIPs, has also been shown to be suitable for the detection of IL-6, as a biomarker for Alzheimer's disease, and the development of POCT devices [[100,](#page-27-15) [106](#page-27-22)]. The MIP film was obtained by the co-electropolymerization of pyrrole and carboxylated pyrrole in the presence of IL-6 protein, and SPCEs. This MIP-based sensor showed a detection limit at a level below pg/mL for IL-6 in spiked serum, good sensitivity, reproducibility, accuracy, and fast response being promising for the development of early diagnostics POCT for neurological diseases [[100](#page-27-15)].

Some relevant and recent examples of electrochemical (bio)sensors for the detection of biomarkers involved in neurological disorders are presented in Table [3](#page-12-0).

#### **COVID‑19‑associated biomarkers**

Viral infection outbreaks are a great menace to groups of individuals due to their fast and easy spread but also for the incubation period which varies depending both on the viral pathogen and the infected person. The contemporary COVID-19 outbreak became in such a short time a matter of great concern for the global population. As of half of June 2022, according to WHO, there were reported almost 532 million confrmed cases and 6 million deaths, respectively [\[108](#page-27-23)].

SARS-CoV-2 disease stands for "severe acute respiratory syndrome coronavirus 2." This virus makes patients experience in the frst instance fu-like symptoms with possible outcomes such as acute respiratory distress syndrome (ARDS) and multiple organ failure (MOF) [\[109,](#page-27-24) [110](#page-27-25)], and the severity of the disease is topped up by an important pathophysiological aspect—the aggressive infammatory response, which causes serious damage to the airways [[109,](#page-27-24) [111](#page-27-26), [112](#page-27-27)].

The SARS-CoV-2 virus has four main structural proteins (Fig. [3A\)](#page-14-0): membrane protein (M), an envelope protein (E), spike protein (S), and one multifunctional structural nucleocapsid protein (ncovNP). The N protein is the main component of the ncovNP and is responsible for packaging and protecting coronavirus the genomic RNA, being divided into five distinct regions (Fig.  $3B$ ) [[12](#page-24-11)]. The S protein is cleaved into two separate polypeptides S1 and S2 within the host by a furin-like protease  $[113]$  $[113]$ . Due to its affinity toward the human angiotensin-converting enzyme 2 (hACE2), it easily penetrates and infects human cells [[114](#page-28-1), [115](#page-28-2)].

<span id="page-12-0"></span>



Table 3 (continued)

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protein, *NFL* neuroflament protein, *AuNFs* Au nanofowers, *L-cys* L-cysteine, *PARK7/DJ-1* protein deglycase DJ-1, *bio-PET* bio-based poly(ethylene terephthalate), *HUVEC* human umbilical estradiol, *EE* ethynylestradiol, *E3* estriol, *AMX* amoxicillin, *LVX* levofoxacin, *AuPd-ANPs/GQDs/CF* AuPd alloy nanoparticle–decorated graphene quantum dots on carbon fber, *Cys* cysteine, CA cysteamine, Hcys homocysteine, NAC-N acetylcysteine, GSH glutathione, DTE 1,4-dithioerythritol, DHLA dihydrolipoic acid, NCNTAs nitrogen-doped carbon nanotube arrays (NCNTAs), FIB fibronectin, TNT troponin-t, FIP fipronil, sTREM-1 soluble triggering receptor expressed on myeloid cells-1, CYFRA-21 cytokeratin 19 fragment, HB hemoglobin, TGF- $\beta$ 1 transforming FIB fibronectin, TNT troponin-t, FIP fipronil, sTREM-1 soluble triggering receptor expressed on myeloid cells-1, CYFRA-21 cytokeratin 19 fragment, HB hemoglobin, TGF-β1 transforming growth factor [31, RANKL receptor activator of nuclear factor kappa-B ligand, CRP C-recative protein, CCL-2 chemokine ligand 2, CeNF cerium oxide nanofiber, MF-4W multifunctional DNA growth factor β1, *RANKL* receptor activator of nuclear factor kappa-Β ligand, *CRP* C-recative protein, *CCL-2* chemokine ligand 2, *CeNF* cerium oxide nanofber, *MF-4WJ* multifunctional DNA 4-way junction, *PS* polystyrene, *PAMAM-NFs* polyamidoamine dendritic polymer nanofbers, *GSPPE* graphene screen–printed paper electrode, *DDAH* diblock dual-aptamer allosteric hairpin, *SFG* graphene composite, *AuHCF* gold hexacyanoferrate, *AB* acetylene black, *EpxS-Ppyr* epoxy-substituted-poly(pyrrole) polymer, *GC* glassy carbon, *SAV-pHRP* streptavidin-poly horseradish protein, NFL neurofilament protein, Au/VFs Au nanoflowers, L-cys L-cysteine, PARK7/DJ-1 protein deglycase DJ-1, bio-PET bio-based poly(ethylene terephthalate), HUVEC human umbilical vein endothelial cells, CVTs carbon nanotubes, PDMS polydimethylsiloxane, NiCo2S4@CoS2 NAs@CC nickel cobalt sulfide/cobalt sulfide nanostructured arrays on carbon cloth, Raw 264.7 vein endothelial cells, *CNTs* carbon nanotubes, *PDMS* polydimethylsiloxane, *NiCo2S4@CoS2 NAs@CC* nickel cobalt sulfde/cobalt sulfde nanostructured arrays on carbon cloth, *Raw 264.7* mammalian cells, CC/Co@C-CNTs Co nanoparticles embedded both in carbon sheets and in the tips of CNTs, CSF cerebrospinal fluid; MDA-MB-231 cells human breast cancer cell lines, E2 CA cysteamine, Hcys homocysteine, NAC-N acetylcysteine, GSH glutathione, DTE 1,4-dithioerythritol, DHLA dihydrolipoic acid, NCNTAs nitrogen-doped carbon nanotube arrays (NCNTAs), 4-way junction, PS polystyrene, PAMAM-NFs polyamidoamine dendritic polymer nanofibers, GSPPE graphene screen-printed paper electrode, DDAH diblock dual-aptamer allosteric hairpin, SFG graphene composite, AuHCF gold hexacyanoferrate, AB acetylene black, EpxS-Ppyr epoxy-substituted-poly(pyrrole) polymer, GC glassy carbon, SAV-pHRP streptavidin-poly horseradish *LSV* linear sweep voltammetry, *AβO* amyloid β oligomer, *CFP/AuPt* superhydrophobic carbon fber paper with AuPt alloy nanoparticles, *Aβ40, Aβ42* amyloid beta oligonucleotides, *tau441* mammalian cells, *CC/Co@C-CNTs* Co nanoparticles embedded both in carbon sheets and in the tips of CNTs, *CSF* cerebrospinal fuid; *MDA-MB-231 cells* human breast cancer cell lines, *E2* Cys cysteine, estradiol, EE ethynylestradiol, E3 estriol, AMX amoxicillin, LVX levofloxacin, AuPd-ANPs/GQDs/CF AuPd alloy nanoparticle-decorated graphene quantum dots on carbon fiber, peroxidase, CMA 4-carboxymethylaniline, EC-MIS electrochemical magnetoimmunosensor peroxidase, *CMA* 4-carboxymethylaniline, *EC-MIS* electrochemical magnetoimmunosensor

Upon the infection with the SARS-CoV-2 virus, the B cells of the host immune system respond and produce antibodies against the N-protein. Antibodies for S-protein are formed only after 4–8 days since the frst symptoms appear. However, the Ig antibodies against SARS-CoV, such as IgA, IgM, and IgG, can be detected at diferent moments after the illness onset: IgA and IgM by the third week and IgG by the sixth week, respectively. Nevertheless, the accuracy of the detection reduces to 84% and 53% for IgG and IgA and IgM, respectively, after the seventh week since the patient got infected  $[116–118]$  $[116–118]$ .

The reference detection method for SARS-CoV-2 infection uses real-time reverse transcriptase polymerase chain reaction (RT-qPCR) for the amplifcation of specifc viral sequences from the virus genome such as E, RdRP, N, and S genes. Besides the requirements for skilled and trained personnel as well as bio-safety level-3 (BSL-3) laboratories, possible mutations of the SARS-CoV-2 virus can reduce the accuracy of the gold detection method. Given these inconveniences, the identifcation of prognostic biomarkers that can forecast the clinical course of a patient and the development of fast and accurate POC assays that provide (i) information on diagnosis, (ii) correlation with the immune response, and (iii) evaluation of the severity of the disease remain important objectives for the scientifc community.

Therefore, this section evaluates possible prognostic markers involved in the immune-pathogenesis of SARS-CoV-2 infection and provides an overview of the POC electrochemical assays developed to confrm the infection with the SARS-CoV-2 virus.

#### **Possible prognostic markers of the immune response**

Several biomolecules involved in hyper-immune responsiveness, bradykinin and cytokine storms, and vascular leakage are assumed to be potential prognostic biomarkers for the COVID-19 disease [[109](#page-27-24), [119,](#page-28-5) [120\]](#page-28-6). A list of potential biomarkers to predict the outcome of the COVID19 disease was realized by Mariappan et al. [\[119\]](#page-28-5) (Table [4](#page-15-0)). The level of some cytokines, including the previously mentioned ones, undergoes a characteristic change in the so-called cytokine storm. It has also been observed that in post-COVID, their level remains elevated [\[119\]](#page-28-5). This may be caused by the appearance of some infammatory impairments located in some organs or tissues. The importance of cytokines in the detection of the COVID infection and the post-COVID status consists both in the rapid identifcation of the infection and in the determination of the general status of the patient and the evaluation of the prognosis. This can be particularly relevant in clinical practice, especially in the case of patients with severe damage and associated comorbidities. However, these need to be frst validated by clinical studies and prove their efficacy in predicting the degree of the disease's severity and long-term efects.



<span id="page-14-0"></span>**Fig. 3** Schematic representation of **A** SARS-CoV-2 virus; **B** domain architecture of SARS-CoV-2 N protein. Reprinted with permission by Elsevier from [\[12\]](#page-24-11)

Type of biomarkers	Representatives
Cytokine storm biomarkers	IL-1, IL-2, IL-6, IL-7, IL-12, IL-17, IL-18, TNF- $\alpha$ , M-CSF, G-CSF, CXCL-10/IP-10, CCL-3, CCL-5, IFN- $\gamma$ , $MCP-1$
Macrophage biomarker	CD14, CD163, TLR2, TLR4, CD86, CD80, CD68, SOCS3, CD200R, CD206, ferritin
Endothelial biomarkers	Endoglin, syndecan-1, endothelin-1, cluadin-5, angiopoietin-1 (Ang-1), Ang-2, PECAM, S1P, VCAM, vWF, Tie2
Bradykinin storm biomarkers	Bradykinin (BK), des-Arg9-bradykinin (DAKB), bradykinin-1 receptor (B1R), B2R, neprilysin (NEP), kal- likrein, kininogen (LW & HW)
Clinical biochemical biomarker	D-mer, C-reactive protein (CRP), ferritin

<span id="page-15-0"></span>**Table 4** List of potential biomarkers to predict the COVID disease short- and long-term outcomes (reprinted with permission by Elsevier from [[119\]](#page-28-5))

*IL-1* interleukin-1, *IL-2* interleukin-2, *IL-6* interleukin-6, *IL-12* interleukin-12, *IL-17* interleukin-17, *IL-18* interleukin-18, *TNF-α* tumor necrosis factor alpha, *M-CSF* macrophage colony-stimulating factor, *G-CSF* granulocyte colony-stimulating factor, *CXCL-10* C-X-C motif chemokine ligand 10, *CCL-3* chemokine (C–C motif) ligand 3, *CCL-5* chemokine (C–C motif) ligand 5, *IFN- γ* interferon gamma, *MCP-1* monocyte chemoattractant protein-1, *CD14* cluster of diferentiation 14, *CD163* cluster of diferentiation 163, *TLR2* Toll-like receptor 2, *TLR4* Toll-like receptor 4, *CD86* cluster of diferentiation 86, *CD80* cluster of diferentiation 80, *CD68* cluster of diferentiation 68, *SOCS3* suppressor of cytokine signaling 3, *CD200R* receptor for membrane glycoprotein OX-2, *CD206* cluster of diferentiation 206 also known as the mannose receptor C type 1 (MRC1), *PECAM* platelet endothelial cell adhesion molecule-1, *S1P* sphingosine-1-phosphate, *VCAM* vascular cell adhesion molecule, *vWF* von Willebrand factor, *Tie2* receptor of angiopoietin

# **POCT electrochemical assays for SARS‑CoV‑2 virus detection**

Besides the challenges of analysis methods encountered when subjected to miniaturization, to enable the production of POCT tools, the quantifcation of SARS-CoV-2 must be realized at low levels and fast pace. Otherwise, false-negative results could increase the spread rate of the disease through incorrectly tested individuals. Compact and costefective devices for fast and accurate detection of the virus are the need of the hour. Recently published POCT electrochemical devices for the detection of COVID19-related diagnostic markers were analyzed and the most important characteristic and fgures of merit are summarized in Table [5.](#page-16-0) For more exhaustive research on COVID19-associated diagnosis tools reporting other detection mechanisms, besides the electrochemical ones, several recent reviews are recommended[[121–](#page-28-7)[126\]](#page-28-8).

Electrochemical immunosensors for the N-protein of SARS-CoV-2 based on lab-produced and purified anti-SARS-CoV-2 ncovNP antibodies were evaluated in clinical samples. For example, Białobrzeska et al. [[12\]](#page-24-11) reported an immunosensor for the RNA binding domain of ncovNP of SARS-CoV-2 at diferent electrodes (diamond/gold/GCE). The highest non-adsorption effect was seen at the BDDEmodifed biosensor, regardless of the low capacity and background currents extending sensory efficiency  $[12]$  $[12]$ . Different strategies to replace biological receptors were investigated in the last decades. Besides aptamers as biomimetic elements, MIPs electrochemically generated have been extensively exploited for the sensitive and selective detection of COVID19-related biomarkers based on the antibody-like afnity that allows discrimination among similar molecules

[[148](#page-29-0)]. A more sensitive electrochemical approach for the detection of SARS-CoV-2 ncovNP was developed by Raziq et al. [\[127](#page-28-9)] based on a MIP disposable sensor chip at thinflm electrodes (TFE). The sensor showed an LOD about twice higher in the nasopharynx swabs of patients (27 fM) [[127\]](#page-28-9), but still about 3 orders of magnitude lower than the immunosensors developed by Białobrzeska et al. [\[12\]](#page-24-11).

A multiplexed, portable, wireless electrochemical platform was developed for the analysis of ncovNP, IgM and IgG antibodies, and the infammatory biomarker C-reactive protein in saliva and blood at an electrochemical device of four working electrodes of laser-engraved graphene (LEG). The graphene surface was modifed by drop-casting the SARS-CoV-2-associated antigen, N-protein, and S1 with the help of a cross-linker reagent to bind with selective immunoglobulin isotypes against both antigens in the same electrode [\[128\]](#page-28-10). Another graphene-based electrochemical approach was realized for the detection of the N-gene of SARS-CoV-2 in nasopharyngeal and nasal swab and saliva within 5 min. However, the affinity was realized by means of the hybridization reaction with an antisense ssDNA strand that was immobilized at a graphene-modifed paper surface, leading to increased sensitivity [[129\]](#page-28-11).

The nucleocapsid and spike proteins of SARS-CoV-2 are the most promising biomarkers for antigen-based rapid diagnostics tests. A sandwich immunosensor enabled the detection of the S-protein with a good sensitivity and a LOD in the ng/mL range, using the streptavidin (poly) HRP-80 enzymatic complex as an amplification system [[130](#page-28-12)]. Another approach that implemented the chronoamperometry for the fast detection (5 min) of S-protein was based on the combination of antibodies against the specifc antigen for increased selectivity and DNA linkers tethered with a

<span id="page-16-0"></span>

Table 5 (continued) **Table 5** (continued)

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IFE gold-based thin-film electrodes, AuNPs gold nanoparticles, BCoV bovine coronavirus, BDDE boron-doped diamond electrode, BSA bovine serum albumin, CA ohronoamperometry, CB tive protein, CV cyclic voltammetry, cTnI cardiac troponin I, CP capture probe, CysOH mercaptoethanol, DPV differential pulse voltammetry, E2 surface viral antigen, EBV Epstein-Barr virus, EBoV Ebola virus, EIS electrochemical impedance spectroscopy, Fc ferrocene, FET field-effect transistor, FTO fluorine-doped tin oxide electrode, GCE glassy carbon electrode, GO graphene NL63 human coronavirus strain NL63, HIV human immunodeficiency virus, HRP horseradish peroxidase, IL-6 interleukin-6, IVA influenza A virus, IVB influenza B virus, InfA-S-gene influenza A spike protein, *JEV* Japanese encephalitis virus, *MB* magnetic beads, *Mb* methylene blue, *MAA* mercaptoacetic acid, *M-S-gene* MERS-CoV spike protein, *NCM* nitrocellulose membrane, *TFE* gold-based thin-flm electrodes, *AuNPs* gold nanoparticles, *BCoV* bovine coronavirus, *BDDE* boron-doped diamond electrode, *BSA* bovine serum albumin, *CA* chronoamperometry, *CB* carbon black, CD48 cluster of differentiation 48 protein, CMV cytomegalovirus, CNF carbon nanofibers, Co-TNTs cobalt-functionalized TiO<sub>2</sub> nanotubes, CP chronopotentiometry, CRP C-reacive protein, CV cyclic voltammetry, cTnI cardiac troponin I, CP capture probe, CysOH mercaptocthanol, DPV differential pulse voltammetry, E2 surface viral antigen, EBV Epstein-Barr virus, EBoV Ebola virus, EIS electrochemical impedance spectroscopy, Fc ferrocene, FET field-effect transistor, FTO fluorine-doped tin oxide electrode, GCE glassy carbon electrode, GO graphene oxide,  $\beta$ -1,4-GALT-5  $\beta$ -1,4-galactosyltransferase 5, HBsAg hepatitis B surface antigen, HCV hepatitis C virus, HCoV human coronavirus, HCoV-229E human coronavirus strain 229E, HCoV-NL63 human coronavirus strain NL63, HIV human immunodeficiency virus, HRP horseradish peroxidase, IL-6 interleukin-6, IVA influenza A virus, IVB influenza B virus, InfA-S-gene influenza A spike protein, JEV Japanese encephalitis virus, MB magnetic beads, Mb methylene blue, MA mercaptoacetic acid, M-S-gene MERS-CoV spike protein, NCM nitrocellulose membrane, ncovNP SARS-CoV-2 nucleoprotein, NoV norovirus, PmPD poly-m-phenylenediamine, Pneu Streptococcus pneumoniae antigen, RBD receptor-binding domain, RCA rolling circle amplification, rGO reduced graphene oxide, RuV recombinant Rubella virus, RSV respiratory syncytial virus, SJ subunit of SARS-CoV-2 spike protein, Si-MOSFET metal-oxide semiconductor fieldeffect transistor, SiNPs silica nanoparticles SCX8 p-sulfocalix[8]arene, SPCE screen-printed carbon electrode, SWV square wave voltammetry, S-gene SARS-CoV-2 spike antigen protein, TMB efect transistor, *SiNPs* silica nanoparticles SCX8 p-sulfocalix[8]arene, *SPCE* screen-printed carbon electrode, *SWV* square wave voltammetry, *S-gene* SARS-CoV-2 spike antigen protein, *TMB* 2-ME 2-mercaptoethanol, DTSSP 3,3'-dithiobis [sulfosuccinimidyl propionate], 4-ATP 4-aminothiophenol, 4-ABA 4-aminobenzoic acid, AO acridine orange, Au-E gold-based electrodes, Au-2.ME 2-mercaptoethanol, DTSSP 3,3'-dithiobis [sulfosuccininidyl propionate], 4-ATP 4-aminothiophenol, 4-ABA 4-aminobenzoic acid, AO acridine orange, Au-E gold-based electrodes, Aucarbon black, *CD48* cluster of diferentiation 48 protein, *CMV* cytomegalovirus, *CNF* carbon nanofbers, *Co-TNTs* cobalt-functionalized TiO2 nanotubes, *CP* chronopotentiometry, *CRP* C-reacoxide, *β-1,4-GALT-5* β-1,4-galactosyltransferase 5, *HBsAg* hepatitis B surface antigen, *HCV* hepatitis C virus, *HCoV* human coronavirus, *HCoV-229E* human coronavirus strain 229E, *HCoVncovNP* SARS-CoV-2 nucleoprotein, NoV norovirus, *PmPD* poly-m-phenylenediamine, *Pneu Streptococcus pneumoniae* antigen, *RBD* receptor-binding domain, *RCA* rolling circle amplifcation, *rGO* reduced graphene oxide, *RuV* recombinant Rubella virus, *RSV* respiratory syncytial virus, *S1* subunit of SARS-CoV-2 spike protein, *Si-MOSFET* metal-oxide semiconductor feld-3,3',5,5'-tetramethylbenzidine, Au-TFME thin-film gold electrode, PAPBA poly(3-aminophenylboronic acid), TB toluidine blue, VSV vesicular stomatitis virus 3,3′,5,5′-tetramethylbenzidine, *Au-TFME* thin-flm gold electrode, *PAPBA* poly(3-aminophenylboronic acid), *TB* toluidine blue, *VSV* vesicular stomatitis virus redox probe (ferrocene) for signal achievement [[131](#page-28-13)]. By contrast, an impedimetric immunoassay enabled the more sensitive detection of S-protein by using AuNPs as electrode modifers [[133](#page-28-15)]. Seo et al. [\[132\]](#page-28-14) reported a graphene-based feld-efect transistor (FET) biosensor to detect S-protein in the fg/mL range in nasopharyngeal swab samples collected from 19 patients. Another FET electrochemical immunosensor allowed the dual detection of S-protein and cardiac troponin I (cTnI) at a gold-modifed glass slide [\[134](#page-28-16)], but the LOD was 100 times higher than DNA-sensor proposed by Seo et al. [\[132](#page-28-14)]. By contrast, Vadlamani et al. [[138\]](#page-28-20) propose the detection of the S-protein within 30 s using a cobaltfunctionalized  $TiO<sub>2</sub>$  nanotube–based sensor. However, both prototypes still need to be validated at diferent levels of S-protein in clinical samples and in the presence of cross-interferents.

Several assays for the detection of the S1 polypeptide of the S-protein were developed based on antibody-target affinity reaction  $[135, 136]$  $[135, 136]$  $[135, 136]$  $[135, 136]$  $[135, 136]$  or biomimetic induced affinity reaction (MIP) [\[137](#page-28-19)]. Although the sensing principles were diferent in terms of the electrode support, detection method, or signal amplifcation, all approaches enabled the detection of the S1 subunit in the fM to pM range.

The simultaneous quantifcation of S- and N-proteins was performed by a sandwich immunoassay that used an enzyme (alkaline phosphatase) for both signal generation and amplifcation. The sensor confguration enabled the detection of S- and N-proteins in untreated saliva as low as ng/mL, within 30 min [[139\]](#page-28-21). One of the recent strategies for DNA amplifcation is represented by the isothermal technique named rolling circle amplifcation (RCA) [[149](#page-29-8)]. A voltammetric biosensor that used the RCA amplifcation for the simultaneous detection of S- and N-genes of SARS-CoV-2 in nasopharyngeal and throat swab was developed by Chaibun et al. [\[140\]](#page-28-22). Silicon nanoparticles were functionalized with acridine orange and MB to target the S- and N-genes, respectively. Sandwich hybridization of redox-active labels with RCA amplicons was realized. The efficiency of the method was proven by the very low LOD of 1.0 copy/mL in clinical samples. Hence, the applicability of the sensor was evaluated in 106 clinical samples, the results being fully in accordance standard RT-PCR method [[140\]](#page-28-22).

Antibody detection from clinical samples has been proven to be a great alternative to the reference RT-PCR method, particularly after the second week of infection. For example, one voltametric [\[141](#page-29-1)] and two impedimetric biosensors were developed for the single detection of SARS-CoV-2 antibodies for the S-protein [[142\]](#page-29-2) and simultaneously with the antibodies for the S1 subunit [\[143\]](#page-29-3). Besides the multiplexed ability of the approach proposed by Ali et al. [\[143](#page-29-3)], the microfuidic channel employed in this sensing strategy enabled the ultrafast (11.5 s) and sensitive (2.8 fM for S1 Ab and 16.9 fM for RBD Ab) detection of the targets. Another advantage of this approach was the regeneration property that was tested in an acidic medium and was performed at least 9 times without any signifcant change in their performance [[143](#page-29-3)]. A very cheap alternative could be represented by electrochemical paper-based devices (ePADs). An ePAD as a screening tool to detect SARS-CoV-2 immunoglobulins (IgG and IgM) was developed by Yakoh et al. [[144](#page-29-4)]. The sensing scheme relies on the disruption of the redox conversion of  $[Fe(CN)_6]^{3-/4-}$  which was triggered by the formation of IgG–S-protein immunocomplex (Fig. [4\)](#page-19-0). The fast (30 min) and sensitive detection of SARS-CoV-2 antibodies was recorded with a LOD of 1 ng/mL that enabled the application of the sensor only in serum samples as for human nasopharyngeal swabs the values of IgG and IgM go down to pg/mL  $[144]$  $[144]$  $[144]$ . Unlike lateral flow immunoassay (LFA) platforms where multiple antibodies are exploited in charge of the reporter, no antibodies are required if label-free electrochemical systems are applied [\[150\]](#page-29-9). Although sensitive and cost-efective, ePADs fail to accurately and uniformly control the fow rate that dramatically infuences the performance of the sensor. To this, capillary-driven microfuidic systems were developed to overcome these limitations [[151](#page-29-10)]. SARS-CoV-2 IgG antibodies for N-protein were detected by a low-cost electrochemical capillary-fow device by chronoamperometry. The antibodies were captured on the nitrocellulose membrane and further electrochemically detected at modifed SPCEs, with a LOD of 5 ng/mL [\[145](#page-29-5)].

A more sensitive electrochemical biosensor for IgG antibodies for N-protein was developed based on the synergistic efect between electrode modifcation with CNF and diazonium chemistry for antigen grafting. Unlike other approaches, the sample collection and detection tools are integrated into the same device by coating the SPCEs with absorbing cotton padding [[146](#page-29-6)]. A sandwich assay electrochemical biosensor for the diagnosis of ORF1ab RNA of SARS-CoV-2 was developed based on a nanocomposite of Au@Fe3O4, GO, p-sulfocalix[8]arene (SCX8), AuNPs, and a fuorescent material, like TB, to obtain Au@SCX8- RGO-TB. The LODs of this assay in synthetic solutions and clinical samples were 3 aM and 200 copies/mL, respectively, being the lowest LOD among the published RNA-based sensors [\[147](#page-29-7)].

As the antibody response is produced only after 4–6 days since the onset of the infection, one must consider the prevalence of false-negative results and the spread of the disease without suitable diagnostic tools at the proper timing.

#### **Sports biomarkers**

Self-monitoring in sports activities has attracted signifcant attention since there are several markers mirroring the physiological and metabolic changes in the human body. Thus,



<span id="page-19-0"></span>**Fig. 4** Schematic representation of the detection procedure of the COVID-19 ePAD for IgG and IgM antibodies. Illustration of the **A** device components, **B** detection principle, and **C** detection procedure of the COVID-19 ePAD. Reprinted with permission by Elsevier [[144\]](#page-29-4)

athletes can monitor their health, performance, and recovery via on-site analysis provided by POC devices. Additionally, dopants can be easily identifed/monitored in performance athletes. Unlike pathological biomarkers, physiological biomarkers tend to vary their levels depending on diet, circadian rhythm, training, and rest cycles. The physiological range of most markers also depends on factors such as age, ethnicity, and gender. For regular monitoring, one must correlate all these variables to fully understand the output data; otherwise, wrong conclusions are drawn. Hence, multiplexed analysis of several biomarkers is of interest for better oversight of the overall performance and health status. Moreover, due to high-tech advancements and continuous improvement of smartphone apps, users and trainers have the possibility to optimize regime training, track their nutrition, and online measure their performances. Wearable sensors are even more popular among athletes, due to attributes such as afordability and ergonomics provided by advances in miniaturized electronics, enabling them to monitor physiological parameters without any expert guidance and during physical activities [\[152](#page-29-11)]. For example, Table [6](#page-20-0) presents a list of wearable electrochemical biosensors for glucose monitoring that can be applied for sport-related activities, but also by patients sufering from diabetes.

More insights about sports biomarker classifcation and their detection by laboratory and POCT can be found in some recent reviews [\[154,](#page-29-12) [155\]](#page-29-13). A list of sports biomarkers was reported and classifed into: *(i) performance markers* (i.e., testosterone, cortisol, growth hormone, insulin-like growth factor-1, erythropoietin, soluble transferrin receptor, hepcidin, lactate, taurine, glutamine, branched chained amino acids, estrogen, progesterone); *(ii) injury, infammation, and recovery markers* (i.e., S-100B, neuroflament light, Tau protein, C-reactive protein, interleukin-6, creatine kinase); and *(iii) nutrition markers* (i.e., glucose, albumin, blood urea nitrogen, vitamin D).

Wearable platform	Sample	Product/company	Current stage	Website
Contact lens	Tears	Smart contact lens, Google, and Novartis	First published in 2014, last update in 2020; the project is on hold	
Smartwatch	Interstitial fluid	GlucoWatch, Cygnus Inc.	FDA-approved, but retracted	$\overline{\phantom{a}}$
Wrist strap	<b>Blood</b>	BioMKR, Prediktor Medical	Under clinical testing for approval and market launch in Europe	https://www.prediktor.com/
Patch		Interstitial fluid Freestyle Libre, Abbott	FDA-approved in the USA in July 2018	https://www.freestyle.abbott/us- en/home.html
Patch	Interstitial fluid	Dexcom G6 CGM, Dexcom	FDA-approved	https://www.dexcom.com/global
Patch	Sweat	MAX3013, Maxim Integrated	Active in production	https://www.maximintegrated. com
Extra-small stick to be placed under the lower eyelid	Tears	NovioSense tear glucose sen- sor, NovioSense	Phase 2 Clinical Trial	http://noviosense.com/

<span id="page-20-0"></span>**Table 6** Wearable electrochemical biosensors for glucose released on the market. Adapted from [\[153\]](#page-29-22)

Injuries among athletes are very common and the determination of the state of the injury is important in the recovery and healing process because it could aid practitioners in implementing the best therapy. Although mainly secreted in high levels into the CSF, the markers are also expressed at lower levels in the bloodstream [[156\]](#page-29-14). To be noted that not many biomarkers reveal the exact state of injuries, the majority of them being non-specifc. Besides injury, infammation is also implicated in the physio-pathological sportscorrelated processes. Biomarkers for muscle damage and infammation often go hand-in-hand leveling up to even 100 times after intensive exercise, being an efect of the body's immune response [[154](#page-29-12)]. Therefore, sensitive POC devices for the detection of injury, infammation, and recovery biomarkers could help athletes to schedule their training sessions depending on their health status.

There are several potential POC devices reporting the analysis of infammation biomarkers, such as cortisol  $[157-159]$  $[157-159]$  $[157-159]$ , C-reactive protein  $[160-162]$  $[160-162]$  $[160-162]$ , Tau protein [\[163](#page-29-19)[–166](#page-29-20)], or S-100B [\[167](#page-30-0)]. For example, cortisol was analyzed using an impedimetric SPCE-modifed electrochemical cell [[157\]](#page-29-15) and a voltammetric Au-interdigitated modifed microelectrode [[158\]](#page-29-21) from saliva and plasma, respectively. By contrast, the detection of cortisol as low as 100 pM in saliva regardless of the presence of interferences such as β-estradiol, testosterone, progesterone, or corticosterone was realized using a voltammetric immunosensor [\[159](#page-29-16)]. Liu et al. [\[167](#page-30-0)] developed a disposable microchip immunosensor for S-100B based on a PMMA channel functionalized with S-100B-specific monoclonal antibodies. Upon the affinity reaction, polyclonal anti-S100 were sequentially coupled by antigen–antibody specifc interaction, followed by the conjugation with alkaline phosphatase (ALP). The strategy enabled the sensitive detection of S-100B as low as 0.1 pg/ mL [[167\]](#page-30-0).

However, these biomarkers are commonly expressed after a traumatic event or in neurodegenerative disorders and a few reported papers in the literature make a connection with the levels over a sports activity. Currently, performance and nutrition markers are more subjected to the development of wearable devices for online monitoring [[155](#page-29-13)].

One challenge is associated with the expression site of infammation markers. Despite performance and nutrition markers, injury and infammation biomarkers are usually measured in serum or plasma. However, infammation may occur locally, in abscesses [\[168](#page-30-1)]. Nevertheless, some latest studies show that there are injury and infammation biomarkers expressed in saliva or perspiration [\[157](#page-29-15), [169](#page-30-2)]; therefore, novel approaches for POC development for applications in sports science could be explored.

# **Conclusions, future perspectives, and challenges**

The improvement of health care services is primarily based on the improvement of the management in the health system, which, in turn, is directly correlated with a quick decision regarding diagnosis and treatment based on a functional diagnostics system, which also benefts from a consistent and well-documented smart therapeutics system. Nowadays, the concept of smart therapeutics must frst of all consider the patient's profle and thus every decision in terms of progression and monitoring evaluation must be personalized with his particularities and needs. The need of the hour is detecting relevant markers at the clinical point-of-care/use by a cheap, user-friendly, sensitive, and reliable detection platform that may facilitate rapid treatment and disease prevention. Especially, medical and sports felds could tremendously beneft from POC-based technologies with superior

specifcity and sensitivity. Hence, fast and reliable analyses may shorten the output of a disease or offer critical information over a patient's health status. However, key points and challenges for POCT accreditation that must be addressed are (i) method performance and validation, (ii) internal and external quality assurance, (iii) staff/user training and competency, and (iv) continuous improvement [[8\]](#page-24-7).

Electrochemical (bio)sensors have attracted attention for the development of smart diagnosis tools with great potential for POCT applications needed for personalized health care, especially due to their advantages, which are related to the suitability for portability and miniaturization, largescale and low-cost production, high sensitivity wide detection range, and low detection limits, as well as easy integration on other systems. The recent developments in the feld of nanoscience and nanotechnology, the discovery of smart electroactive functionalized nanostructures allowing for signal amplifcation, emerging micro-electronics such as miniaturized analyzer, and innovative fabrication technologies such as interdigitated electrodes have also been useful to design and develop miniaturized, automated, and precise sensing systems suitable for clinical applications.

The WHO regulation regarding POCT devices is clear, and according to it, only devices that fully meet the following criteria: "Affordable," "Sensitive," "Specific," "User-friendly," "Robust and rapid," "Equipment-free," and "Deliverable" to all people needing, which are included under the generic name ASSURED, can be included in this category. After a careful analysis of the recent literature data on the subject of this study, there are many studies that meet most of the criteria mentioned in the WHO regulations, these being centralized in Table [7.](#page-22-0) Taking into account that none of the presented devices are commercially available and are not in clinical trials, we included in the "Fully" category all the devices that meet the ASSURED criteria except for the last one, namely to be "Deliverable to all people need." Also, it can be observed that most of the detection devices presented do not meet the "Equipment-free" criterion, and this shortcoming is due to the operation mode of the electrochemical sensors and biosensors, which require the presence of a potentiostat and implicitly an energy source. In several studies from the examples presented, it was possible to solve this shortcoming by developing miniaturized, portable, and ultraportable potentiostats, accessible in terms of price and which allow the operation of the elaborated sensors in a decentralized manner [[36](#page-25-2), [42,](#page-25-17) [71,](#page-26-2) [98](#page-27-14), [104](#page-27-20), [128](#page-28-10)[–130,](#page-28-12) [136,](#page-28-18) [137](#page-28-19), [139](#page-28-21), [141,](#page-29-1) [144](#page-29-4), [146](#page-29-6), [147,](#page-29-7) [149,](#page-29-8) [152](#page-29-11), [158,](#page-29-21) [159\]](#page-29-16). In addition, taking into account the special progress in the feld of electronics in recent years, we are confdent that this shortcoming will be overcome very soon, especially if the advantages that accompany electrochemical detection systems are considered. Among these, we mention the high sensitivity ensured by the easy functionalization with nanomaterials,

the high selectivity when using biological or biomimetic elements, the fast response, but especially their suitability for miniaturization and automation, basic properties in the feld of POCT sensing devices. It can be observed that only one example of detection devices does not meet the criteria "Specific" [\[36](#page-25-2)], respectively, "Sensitive" [[150](#page-29-9)], which can be a justifcation for the large number of recent studies that include electrochemical sensors and biosensors aimed at the development of POCT devices for biomedical and clinical practice, where real, biological samples are very complex in composition and usually contain very low concentrations of analytes.

The development of electrochemical (bio)sensors usually involves a laborious protocol with functionalization steps with nanomaterials and biocomponents, which can represent a drawback. However, the development of the electrochemical sensors is performed before use depending on their stability. Thus, at the moment of the analysis, they can be directly used, obtaining the results easier and faster than in the case of the standard methods applied in clinical analysis. For example, the RT-PCR results may take 1–2 days to report and are expensive, and require laboratory equipment and highly qualifed personnel, rapid POCT diagnostic tests provide the possibility of early detection of infectious COVID-19 cases, and many have been developed since the onset of the disease [[170](#page-30-3)]. Another disadvantage encountered in some of the examples of POCT discussed previously is the lack of comparison with the standard methods used for the analysis of those biomarkers in clinical practice. However, it has been observed recently that more importance has been given to the validation of new methods and devices in relation to standard methods.

Concerning the samples for POC analysis, saliva is considered one of the most compliant biological fuids due to the minimally invasive prelevation methods and ease of manipulation and storage. Being rich in compounds that can be used as markers, saliva is an important tool to assure an easy and simple diagnosis of several diseases [\[119,](#page-28-5) [171](#page-30-4)]. However, further clinical trials are needed to establish the role of each salivary molecule as a marker for health status or disease monitoring as little is known about the correlation between the level of biomarkers in the blood or the fuid where it is secreted and its level in saliva.

Another advantage of using electrochemical sensors is that both potentiostats and electrochemical cells could be integrated into a whole disposable system that would eliminate the need for any dedicated instrumentation as the tests could be run from a smartphone only, making it useable at the point-of-need/use. Portable nano-devices incorporating a cocktail of tests would be ideal for large screening of the population, especially during disease outbreaks, but also for telemedicine diagnosis and monitoring, and sports science. The need for widespread analysis and control over

<span id="page-22-0"></span>



#### Electrochemical sensor configuration/biomarker Meeting the ASSURED criteria Unfulflled criteria Tested in clinical-related settings Commercial availability Ref Electrochemical sensor/SARS-CoV-2 antibodies for spike S1 protein and RBD Fully Deliverable No No [\[144\]](#page-29-4) Paper-based electrochemical sensor/SARS-CoV-2 IgG and IgM for S protein Partially Sensitive; deliverable Yes No [\[150\]](#page-29-9) Electrochemical capillary-fow device/SARS-CoV-2 IgG antibody for N protein Fully Deliverable Yes No [\[146\]](#page-29-6) Cotton-tipped electrochemical immunosensor/SARS-CoV-2 IgG antibody for N protein Fully Deliverable Yes No [\[147\]](#page-29-7) Ultrasensitive sandwich-type electrochemical sensor/ ORF1ab RNA Fully Deliverable Yes No [\[152\]](#page-29-11) Portable and microfluidic electrochemical device/cortisol Fully Deliverable Yes No [\[158\]](#page-29-21) Portable and microfluidic electrochemical device/cortisol Fully Deliverable Yes No [\[159\]](#page-29-16) Disposable electrochemical biosensor/creatine kinase Partially Equipment-free; deliverable Yes No [\[161\]](#page-29-23)

the spread of diferent pathogens has previously faced challenges either due to lack of instrumentation or reduced accuracy of the tests. The use of POCT-based detection devices is more and more often associated with the so-called mobile health (mHealth) equipment. Many wearable sensors (wrist bands, chest straps, and skin mountings) incorporate the integrated sensing transducer, circuitry, biomolecules, and power sources for continuous monitoring of relevant markers. However, the fexibility of materials and users' comfort could sufer some improvements to make wearable sensors more user-friendly and cost-effective [[172,](#page-30-5) [173\]](#page-30-6).

**Table 7** (continued)

Driven by a highly competitive market, innovative technologies, AI, and IoT have become the pile point of companies. Smartphones are beyond being a luxury, but an integral part of human lifestyle, keeping all of us connected to worldwide data. Sports sciences have benefted the most from the smartphone technology, where sports activities and health apps are popular among athletes and smartphone users [\[174\]](#page-30-7). Recent advancements in POCT domain enabled the integration of an analytical platform with smartphones to allow increased mobility, accuracy, and user-friendliness. Although sometimes futuristic, modern technologies have as ultimate goal the development of fast diagnostic and monitoring tools to reduce hospital stays and overall costs and improve life expectancy and style.

Without the aim to cover all POCT electrochemical sensing approaches for the determination of infammationrelated biomarkers, this review treats in a critical manner the importance of this emerging scientifc topic. Nowadays, it is expected that the monitoring of biomarkers involved in infammatory processes occurring in several pathological conditions will gain more interest in healthcare management.

Monitoring infammation biomarkers may aid physicians in their practice. In this regard, the use of various electrochemical approaches toward multiplexed detection of infammatory biomarkers, in particular, can be of much help. Since POCT devices are characterized by small dimensions, low cost, and high sensitivity and it allows the real-time determination of biomarkers through online-assisted equipment, they offer an excellent perspective for easily obtaining an accurate diagnosis. We can assume that all the efforts and research in this feld will overcome the present obstacles and barriers, and the integration of innovative nanomaterials, nanocomposites, microfuidic systems, wireless transmission of the data, and mobile-based biosensing assays will contribute to enhanced overall analytical performance of the developed POCT.

**Author contribution** All authors contributed to the study conception and design. Material preparation and data collection were performed by Diana-Gabriela Macovei and Maria-Bianca Irimes. Data collection and analysis and the frst draft of the manuscript were performed and written by Diana-Gabriela Macovei, Maria-Bianca Irimes, Oana Hosu, and Mihaela Tertis. Cecilia Cristea critically revised the manuscript and supervised the work. All authors reviewed and approved the fnal manuscript.

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# **Declarations**

**Conflict of interest** The authors declare no competing interests.

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