Progress of Polyaniline Glucose Sensors for Diabetes Mellitus Management Utilizing Enzymatic and Non-Enzymatic Detection

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Abstract: Glucose measurement is a fundamental tool in the daily care of Diabetes Mellitus (DM) patients and healthcare professionals. While there is an established market for glucose sensors, the rising number of DM cases has promoted intensive research to provide accurate systems for glucose monitoring. Polyaniline (PAni) is a conductive polymer with a linear conjugated backbone with sequences of single C–C and double C=C bonds. This unique structure produces attractive features for the design of sensing systems such as conductivity, biocompatibility, environmental stability, tunable electrochemical properties, and antibacterial activity. PAni-based glucose sensors (PBGS) were actively developed in past years, using either enzymatic or non-enzymatic principles. In these devices, PAni played roles as a conductive material for electron transfer, biocompatible matrix for enzymatic immobilization, or sensitive layer for detection. In this review, we covered the development of PBGS from 2015 to the present, and it is not even exhaustive; it provides an overview of advances and achievements for enzymatic and non-enzymatic PBGS for self-monitoring and continuous blood glucose monitoring. Additionally, the limitations of PBGS to advance into robust and stable technology and the challenges associated with their implementation are presented and discussed.

Keywords: glucose monitoring; polyaniline; diabetes mellitus; enzymatic; non-enzymatic

1. Introduction

Diabetes Mellitus (DM) is a metabolic disorder characterized by the presence of hyperglycemia due to defects in insulin secretion, insulin action, or both [1]. The complications of DM include nephropathy, neuropathy, retinopathy, foot damage, skin conditions, increased risk of cardiovascular disease, and dementia, to name a few. Moreover, DM can be classified into four categories [2]:

1. Type 1 diabetes (T1DM): Related to autoimmune β-cell destruction, commonly leading to absolute insulin deficiency.
2. Type 2 diabetes (T2DM): Caused by a progressive loss of adequate β-cell insulin secretion, usually on the background of insulin resistance.
3. Gestational DM: High glucose levels diagnosed in the second or third trimester of pregnancy without having a prior diabetic history.
4. Other specific types of diabetes, e.g., monogenic diabetes syndromes, diseases of the exocrine pancreas, and drug- or chemical-induced diabetes.

Currently, 537 million working-age people (20–79 years) have diagnosed or undiagnosed DM. This number is projected to increase to 643 million and then to 783 million
by 2030 and 2045, respectively [3]. In addition, the most significant increase will occur in regions where economies move from low to middle income status. The alarming increase in the worldwide prevalence of DM has increased interest in developing easy-to-use devices to support diagnosis and treatment. These devices require accurate self-assessment of glucose, the so-called golden standard for DM monitoring. Specifically, continuous glucose monitoring is a tool that helps significantly to prevent the risks of other diabetes-induced comorbidities, allowing people to maintain a healthy lifestyle, avoiding costly and lethal late-stage diabetic complications [4,5]. In 1962, Clark and Lyons manufactured the first enzymatic glucose sensor, which established the basis of self-assessment glucose monitoring. The fundamental principle is based on the control of oxygen consumption as a function of the catalytic oxidation of glucose by glucose oxidase (GO\(_x\)) enzyme [6]. While the sensor presented several drawbacks, this breakthrough advance evolved into a global, successful, and well-established market. Since then, several innovative configurations and materials have been incorporated for glucose sensors including, metal nanoparticles [7], carbon-based nanomaterials [8,9], metallic oxide [10], and conductive polymers (CPs) [11–13].

CPs such as polypyrrole (PPy), polythiophene, poly(3,4-ethylenedioxythiophene) (PEDOT) have been incorporated in novel glucose sensors; however, PANi has been extensively applied for chemical sensors and is closely related to the glucose sensing field. The application of PANi in chemical sensing arose in the mid-1980s when the first sensor for pH monitoring appeared [4]. At the end of this decade, the first studies of PANi-based glucose sensors (PBGS) also emerged, with Shinohara et al. reporting platinum electrodes and microelectrodes modified with electropolymerized PANi in the presence of GO\(_x\). In this case, the PANi allows enzyme stability and functions as a charge transfer interface between the enzyme and the electrode [5]. Figure 1 presents a graphical description of PBGS evolution, encouraged by CPs’ Nobel award, and formally started with Shinohara sensor device. Since then, there has been an increasing trend for the development of PBGS with novel morphologies and assemblies using PANi as base material. In this review, we covered the development of PBGS from 2015 to the present, using enzymatic and non-enzymatic mechanisms. We first introduced the principles of glucose mechanisms in both configurations, then we presented a background of the main PANi characteristics and their suitability for glucose detection. Subsequently, we collected references for enzymatic and non-enzymatic PBGS, highlighting the main achievements of these devices for glucose detection intended for DM care.

Figure 1. Graphical description of PANi-based glucose sensors (PBGS) evolution.

2. Glucose Sensing: Mechanisms for Enzymatic and Non-Enzymatic Detection
2.1. Mechanisms for Glucose Detection

The GO\(_x\) enzyme has been the most widely employed in enzymatic biosensors due to its relatively high selectivity, sensitivity, and stability. The main component of the large protein molecule is the redox center, flavin adenine dinucleotide (FAD). This redox center is located inside the enzyme, protected by a thick layer of protein. The FAD is reduced by interacting with glucose to dihydro flavin adenine dinucleotide (FADH\(_2\)), producing the subproduct gluconate (Figure 2). The electron transfer to the active center is a major limiting factor and the cause of complicated electron transfer mechanisms. There are currently two approaches for oxidizing the reduced FADH\(_2\) center: Catalytic regeneration, reaction with a mediator, and direct electron transfer to the electrode (DET). Therefore, the glucose
biosensor based on GO\textsubscript{x} has evolved in three generations, and a fourth has come along without involving a biomolecule. In the following sections, each strategy will be discussed.

\[
\text{GO}_x(\text{FAD}) + \text{Glucose} \rightarrow \text{GO}_x(\text{FADH}_2) + \text{Glutonate}
\]

**Figure 2.** Mechanism for oxidation of glucose by glucose oxidase (GO\textsubscript{x}) enzyme.

### 2.1.1. First-Generation Biosensors

The first-generation of glucose biosensor is based on the catalytic regeneration of the FAD center in the presence of oxygen as a co-substrate. The first oxygen electrode was developed by Clark and Lyon [6], using GO\textsubscript{x} coated by a semipermeable dialysis membrane on an electrode.

The dependence on free oxygen as a catalytic mediator and electroactive interference species in the blood are two main problems this biosensor faces. For instance, there is a low concentration of oxygen in blood samples, which is not enough to maintain glucose oxidation efficiently; so, this oxygen deficit has a significant impact on the accurate determination of glucose levels. The associated mechanism for first-generation enzymatic biosensors is shown in Figure 3.

\[
\text{GO}_x(\text{FADH}_2) + \text{O}_2 \rightarrow \text{GO}_x(\text{FAD}) + \text{H}_2\text{O}_2
\]

**Figure 3.** Reduction of GO\textsubscript{x} (FADH\textsubscript{2}) by O\textsubscript{2} for first-generation glucose biosensors.

### 2.1.2. Second-Generation Biosensors

Due to the oxygen dependence problems and electroactive interferences observed in first-generation enzymatic glucose sensors, alternative co-substrates were used, namely mediators. Furthermore, the synthetic electron acceptance mediators make it easier to transfer electrons from the enzyme’s redox-active site to the conductive electrode and subsequent reoxidation by the electrode, resulting in a quantifiable current. Several non-physiological mediators have been studied, including those derived from ferrocene [14] and ferricyanide [15], quinones [16], and transition metal complexes [17].

Moreover, the mediators generally have insolubility and low molecular weight characteristics, making them ideal for enzymatic glucose analysis. In addition, mediators possess reversible or quasi-reversible properties, a lower redox potential to avoid the oxidation of the interfering species, high stability, resistance to the formation of secondary compounds, and low toxicity. Nevertheless, it is complicated to maintain contact between species (enzyme-mediator-electrode) because the mediators are small and more diffuse during relatively prolonged device use [15,16].

Another drawback is that the mediator reacts with the enzyme at a rate considerably faster than oxygen. However, it causes a competitive reaction between the dissolved oxygen-enzyme and the mediator-enzyme, thus reducing the efficiency of the system and causing an accumulation of hydrogen peroxide.
Over the years, a variety of organic and inorganic compounds have been studied as electron transfer mediators. For instance, osmium complexes [18,19], ferricyanide [20,21], and their derivatives meet the mainly required criteria for an excellent mediator. These requirements are low operational potential, high chemical content, stability of both redox forms, rapid reaction with reduced enzyme (to minimize O$_2$ competition), and low solubility in aqueous media.

In recent years, the enzymatic glucose sensor developed by Heller et al. [22] for Abbott Diabetes Care is a clear example of this second generation [23]. The FreeStyle blood is a typical example of the second-biosensors generations because it employs the glucose dehydrogenase (GDH) enzyme instead of GO$_x$, and osmium complex as mediator. In addition, the measurement method of the glucose monitoring system is microcoulometric and requires a small blood volume (0.3–4.0 µL).

2.1.3. Third-Generation Biosensors

DET is the principle employed in the third generation of enzymatic glucose sensors. The DET from glucose to electrode surfaces through the enzyme's redox center is a challenge that has long been pursued. However, difficulties arise from the thick layer of insulating protein that surrounds the enzyme’s active center that hinder the efficient transfer of electrons with the surface of the electrode. Specifically, for GO$_x$, the FAD/FADH$_2$ center is located at a depth of about 13 Å within the apoenzyme. The advantages of the DET mechanism include removing artificial leachable mediators and operating in a potential window close to the redox potential of the enzyme.

Moreover, glucose and molecules with DET characteristics access this center through the diffusion and penetration of the 3D molecular network. Since this is not possible in a flat electrode, the inclusion of nano and porous materials has raised interest in increasing the surface area and electrode performance [24]. For example, electrodes with mesoporous materials exhibited a surface that traps and encompasses the enzyme; so, a DET from the enzyme to the electrode can occur, and a current related to the oxidation of the enzyme can be registered, without mediators, concerning the overlapping of electroactive interferences, or relying on dissolved oxygen. Nevertheless, the adverse effects of dissolved oxygen competing with the electrode to regenerate the enzyme have not yet been resolved. For this reason, the development of DET-based biosensors continues in the research stage and commercial glucose biosensors employ mainly first and second-generation electrochemical sensors [25].

2.1.4. Non-Enzymatic Glucose Sensors (Fourth Generation)

The interest in non-enzymatic glucose sensors has grown due to the drawbacks of enzyme-based biosensors, such as the possible decrease in the catalytic activity of the enzyme derived from the immobilization process, limited reproducibility, and low stability during prolonged operation. The direct electrocatalytic oxidation of glucose occurs on the electrode surface to avoid deficiencies that originate in the nature of enzymes. A wide variety of materials have been utilized in non-enzymatic glucose sensors, e.g., CPs [11,12], multiwalled carbon nanotubes (MWCNT) [9,13], and molecularly imprinted polymers (MIPs) [26,27], which can mimic enzymes by creating crosslinked polymeric active sites for specific analytes.

The mechanism of non-enzymatic glucose oxidation on the electrode surface is not fully understood and remains an active area of research. However, two main models explain the principles: The first one was proposed by Pletcher [28] and is also known as the activated chemisorption model. In this, the oxidation is started by the adsorption of glucose on the electrode surface. In aqueous solutions, the two most common isomers are $\alpha$-D-glucose and $\beta$-D-glucose, both in their cyclic hemiacetal glucose form. Immediately, the glucose molecule forms a physics interaction with the electrode surface, leading to glucose–metal interaction. Simultaneously, the hydrogen atom located in the hemiacetal carbon is removed from the glucose molecule and attaches to the electrode surface at a site
adjacent to the glucose; this step determines the rate in most electrooxidation processes. Hence, the complex glucose–metal suffered desorption of the glucose molecule because of a change in the oxidation state of the glucose molecule.

Therefore, the formation of intermediate strength is ideal for the adsorption and desorption process to occur. In fact, the proposal of active transition metal centers through the electrode does not consider the oxidative role of hydroxyl radicals that appear absorbed during the electrooxidation of glucose and other organic molecules [29]. A schematic proposal of this model is shown in Figure 4.

Figure 4. Illustration of the glucose–electrode interaction in the Pletcher model.

The second model is known as the “Incipient Hydrous Adatom Mediator” (IHOAM) model, Figure 5, and it is based on the active metal atoms at the electrode surface, which have low reticular stabilization and enhance reactivity. These metal atoms present an oxidation stage prior to the monolayer, during which an incipient hydrous oxide premonolayer (OH\textsubscript{ads}) is formed, which is suggested to mediate the oxidation of glucose on the surface of the electrode. While models can be used to explain glucose oxidation for different electrode materials, both models are still under study [30].

Figure 5. Illustration of the glucose-electrode according to incipient hydrous adatom mediator (IHOAM) model.

3. Polyaniline for Glucose Sensors

3.1. PAni Structure

The application of CPs in the design of biosensors has been reviewed in detail on several occasions in the past [31,32]. Thanks to the ability to transfer electrons produced by biochemical processes, CPs offer significant advantages since they are sensitive to small amounts of the analyte [33]. PAni is a semi-flexible CP of most significant interest for biosensor design due to its electrochemical properties, e.g., reversible redox behavior and electrochemical tuning, simple processability, and long-term stability. Leucoemeraldine, emeraldine salt, and pernigraniline are the three fundamental oxidation states of PAni. Furthermore, it is a versatile material that can have diverse applications in essential areas of science and technology, including electrochromic devices, solar cells, actuators, tissue
engineering, and biosensors. The basic PAni structures are shown in Figure 6, along with their doped and undoped forms.

![Diagram of PAni structures](image)

**Figure 6.** Basic polyaniline structures: Leucoemeraldine, Emeraldine salt, and Pernigraniline.

### 3.2. Conductivity

In CPs, the oxidation state is decisive in their electrical properties; however, PAni has characteristics that differentiate it from the rest. In PAni, electrical conductivity depends on both oxidation state and doping. When PAni in the form of emeraldine base is exposed to a protonic acid (proton doping), nitrogen doping of the imine groups occurs, from which polaron delocalization occurs along the chain, obtaining the conductive form of PAni (emeraldine salt) [34,35]. Proton doping of PAni does not involve the loss or gain of electrons, which occurs in a redox process [36]. Such doping can cause an increase of up to 10 orders of magnitude in the electrical conductivity of PAni [37]. In CPs, the role of the electrons-hole pair is a determining factor in electrical conduction [38,39]. Compared with conventional metals, the CPs do not have a range of energies that are considered unavailable for electrons because the valence and conduction bands overlap. Hence, the band gap or energy gap ($E_g$) in metals is 0 eV. Regarding conductive PAni, the energy gap is ~3.0 eV [40]. Under these conditions, electrons can increase their energy and detach themselves from their atoms by jumping to a higher energy level in the conduction band. For emeraldine base (non-conductive), the $E_g$ between the valence and conduction bands is too large, greater than 3 eV; consequently, the electron is not capable of making such a jump that allows it to ascend to the conduction band, showing poor electrical conduction. It should be noted that experimental and theoretical studies, despite some differences [41], propose that the electronic transport of emeraldine salt is based on polaronic and bipolaronic structures; however, even today, there is no complete consensus [42]. Through doping with
a Brønsted acid, such as hydrochloric acid (HCl), the imine groups are protonated to form the bipolaron. The bipolaron then dissociates to form two polarons, which subsequently rearrange, forming a delocalized polaron in the form of a polysemiquinone (radical-cation salt) [43,44].

3.3. PAni as Matrix Enzyme Immobilization

The factors that influence the enzyme stability are polymerization conditions, incorporation of the enzyme into the matrix of the CP, and dispersion or solution of the enzyme. In particular, concerning PAni-based enzymatic biosensors, it is necessary to consider the polymerization conditions of PAni because emeraldine salt is obtained at pHs lower than 3 (highly conductive). Under these conditions, and in the presence of the polymerization by-products (traces of aniline salt, ammonium persulfate (APS)), enzymes can lose their catalytic activity. On the other hand, the method of incorporation of the enzyme into the CP can also affect the enzyme stability and, therefore, the operation of the sensor. For example, Mu and Xue [45] compared the performance of GOx in electrochemical glucose sensors, as it was supported on the PAni by physical absorption or by in situ polymerization in the presence of the enzyme. The remarkably superior performance of the sensor with the electrochemically deposited enzyme was attributed to the electrochemical doping because the enzyme is strongly bound to the CP through ionic bonds and the desorption towards the electrolyte is lower than when the enzyme is only physically deposited [45]. Other strategies have also been applied to maintain enzyme activity. In this sense, Yan et al. reported an electrode modified with an Au nanoparticles (NPs)–AgCl–PAni biocompatible substrate, on which GOx was immobilized. Electrochemical studies using pH 5 to 9 PBS showed excellent enzymatic activity at pH 6, which is generally adverse for PAni [46]. Subsequently, Zhao et al. reported the modification of an electrode with PAni nanofibers (NFs) synthesized via oxidative polymerization in an interfacial system. GOx was covalently bonded to the NFs through amidation reactions. Circular dichroic and cyclic voltammetry (CV) measurements showed that the covalent immobilization did not affect enzymatic activity [47]. Finally, the use of mixtures of organic solvents with water has also been a tactic used to achieve enzyme stability, e.g., GOx, in the sensor [48], as reported by Lukachova et al., who used water–ethanol mixtures with high ethanol contents. The enzymatic stability in these mixtures and in the presence of a Nafion emulsion as a solid electrolyte was higher than in pure water because the enzyme loses its solubility while remaining in suspension. By this procedure, the enzymatic stability of GOx can be improved compared to traditional methods using highly dilute electrolytes [49]. The subject of enzymatic glucose sensors is discussed more fully later.

3.4. Biocompatibility for Implantable Devices

The requirements of continuous in vivo glucose monitoring in patients with critical conditions of DM have stimulated the development of implantable sensors. Since its inception, the biocompatibility or tolerance of body fluids or tissues with the materials that make up the biosensor has been an aspect of the in-depth study [50]. In addition to interfacial biocompatibility, it is also desirable that body fluids do not cause surface obstruction of the sensor to develop continuous sensing for long periods [51]. It is vital that the patient is not exposed to any danger due to some reaction derived from the sensor implantation, e.g., blood clotting or inflammatory response, because some sensor surface characteristics such as surface粗糙ness and chemical factors such as wettability and surface charge can stimulate poor hemocompatibility that can cause such adverse conditions [52]. CPs also have excellent biocompatibility, as demonstrated by Sun et al. for PAni hydrogels, whose harmlessness towards rat endothelial progenitor cells was demonstrated. In addition to non-toxicity, the PAni hydrogel showed improved cell adhesion and a faster cell proliferation rate [53]. Similarly, Humpolicek et al. [54] conducted biocompatibility studies of PAni, in its conductive and non-conductive forms, with human cell lines. Studies indicated that none of the forms of PAni caused skin damage; however, the emeraldine salt form was
found to exhibit high cytotoxicity. The subsequent purification of the materials showed a considerable reduction in cellular incompatibility, which indicated that the by-products of polymerization are responsible for this effect [54]. Another study conducted in more detail by Kašpárková et al. confirmed that residues of APS, aniline, or aniline salt are highly cytotoxic; therefore, the purification of PANi is of utmost importance towards biomedical applications [55].

4. Polyaniline Based-Enzymatic Glucose Devices

4.1. PANi-Based Glucose Biosensors

Accurate glucose monitoring is a crucial aspect of proper DM management, with enzymatic sensors playing a prominent role in this task. Enzymatic glucose sensors rely on a redox reaction performed on the surface of the electrode driven by the GOx enzyme. The commonly found electrode materials include glassy carbon electrodes (GCE) [56,57] screen-printed carbon electrodes (SPCE) [58,59] and Platinum (Pt) electrodes [60,61]. In PBGS, these supports are subsequently modified with PANi obtained by a variety of methodologies such as chemical oxidative polymerization (COP) [56], electrochemical polymerization (ECP) [62], free radical polymerization [59], plasma polymerization [63] and interfacial polymerization [64]. The sensing mechanisms of PBGS have followed the innovation path described in Section 2.1, acting as a transducer for detecting either O₂ or H₂O₂ subproducts in the first generation, as electron mediator in the second generation, and even as a molecular wire for DET in the third generation. Figure 7 shows the mechanism of detection followed by electrochemical PBGS in all generations.

![Figure 7. Electrochemical PBGS: First generation, second generation, third generation and non-enzymatic.](image)

A key aspect for optimal performance of electrochemical biosensors is the proper immobilization of enzymes over the electrode, which can be achieved through physical and chemical strategies [65]. When PANi is incorporated over the working electrode, the modified surface exhibits suitable characteristics for enzyme immobilization such as enhanced conductivity, high surface area, available functional groups for linkage and biocompatibility. The most straightforward immobilization technique for PBGS is physical absorption of the GOx through electrostatic attraction and Vander Walls forces; however, the poor long-term operational stability is a major drawback of this technique [66]. The immobilization by crosslinking with glutaraldehyde or covalent bonding through the reaction with amino (NH₂) functional groups allows high enzyme loading and enhances operational stability. Nevertheless, the chemical reactions can lead to structural changes of the enzyme [67]. A convenient and highly reported immobilization method is the direct entrapment of the enzyme during the ECP of PANi. In this one-step preparation method, the aniline monomer and GOx are in the same solution, producing biofilms on the electrode surface by controlling applied potential, time, and enzyme amount [65].
representation of the main steps of the immobilization strategies for GOx using PANi is presented in Figure 8.

![Figure 8. Immobilization strategies for GOx in PANi-based biosensors: (a) Physical absorption, (b) covalent bonding, (c) crosslinking, (d) entrapment by ECP.](image)

In addition to the immobilization matrix, a prominent role of PANi is as electrode material for electrochemical transduction. Enzymatic glucose biosensors with bare PANi as transducer take advantage of the polymer key features such as high conductivity, inherent electroactivity and adequate support for electronic transfer. Electrodes modified with sole PANi can exhibit different morphologies such as microtubes [68], nanofibers, nanoflakes [69], and nanoflowers [70], acting as transduction elements and immobilization matrix for GOx at the same time. However, the results for PANi as a single electrode element still present challenges, including limited detection range and low operational stability in the long term. To improve these issues, copolymerization has been proposed for tuning specific characteristics of PANi, e.g., conductivity [64] or solubility [71] of the polymer. For example, to improve the processability of PANi, Tugrul Cem Biak developed a PANi-poly(ethylene glycol) graft copolymer (PANi-g-PEG) combining COP and click chemistry. The resulting water-soluble copolymer allowed a simpler method for GOx immobilization through electrostatic interaction and hydrogen bonding, simplifying the biosensor manufacturing process [71].

However, for developing PBGS, the most widely reported strategy is the assembly of ternary and quaternary composites with a plethora of materials, including carbon-based nanomaterials (e.g., graphene [72,73], MWCNT [57,74]), metallic nanoparticles (e.g., Au [57,75], Pt [76,77]), and polymers (e.g., PEO [78], poly(acrylic acid) [79], and SDS [80]). An example of these strategies was presented by Maity et al., developing a composite of aminated (NH$_2$) MWCNT/PANi/rGO/AuNPs over an SPCE [58]. PANi was synthesized by COP and mixed by sonication with NH$_2$-MWCNT, rGO, and AuNPs. The obtained nanocomposite exhibited convenient characteristics such as high surface
area, porosity, and available sites for GOx covalent immobilization (Figure 9A). In this case, PANi acted as a conductive binder of the assembly over the surface of SPCE. The NH2-MWCNT/PAni/rGO/AuNPs/GOx biosensor showed good electrocatalytic activity, following the detection mechanism described by first glucose biosensor generation. The obtained linear detection range was 1 to 10 mM, a sensitivity of 246 µA mM⁻¹ cm⁻², and a limit of detection (LOD) of 63 µM. Additionally, the biosensor performed well in the presence of interferences such as dopamine, uric acid, and ascorbic acid. The device was evaluated in blood serum samples for practical applications, achieving a recovery rate above 96.5% and retaining 93.4% of the initial activity after 60 days, when stored at −20 °C.

Figure 9. Applications of PANi in enzymatic biosensors: (A) Preparation of NH2-MWCNT/PAni/rGO/AuNPs/GOx first generation biosensor: (a) synthesis of NH2-MWCNT, (b) rGO, (c) COP of PANi, (d) synthesis of AuNPs, (e) assembly of NH2-MWCNT/PANI/rGO/AuNPs for GOx immobilization; (B) the assembly of PAA/rGO/VS-PANI/LuPC2/GOx/MHF for second generation biosensor (a), a view of the obtained MFH (b) along with SEM images of the morphology of PAA/rGO/VS-PANI/MHF (c) and MFH morphology without rGO (d); (C) synthesis of PANi/TiO2NT composite for third generation biosensor (a), and SEM images of PANi (b) and TiO2NT (c). Figures reproduced with permission from Debasis Maity et al., H. Al-Sagur et al., and Jie Zhu et al. [58,59,81].
In a different strategy, H. Al-Sagur et al. reported a three-dimensional hydrogel assembled with polyacrylic acid (PAA), aminated RGO, vinyl substituted PANi (VS-PAni), and lutetium Phthalocyanine (LuPc$_2$), Figure 9B [59]. The so-called multifunctional hydrogel (MFH) was obtained using free radical polymerization by incorporating VS-PAni with rGO and PAA during the synthesis. Then, the MFH was cast over an SPCE and subsequently doped with LuPc$_2$ to be applied as support for GO.$x$. The characterization revealed a hierarchical structure with low charge transfer resistance and interconnected networks of VS-PAni. The proposed mechanism corresponds to the second generation, with LuPc$_2$ acting as a transfer mediator. The PAA/rGO/VS-PAni/LuPc$_2$/GO.$x$/MHF biosensor resulted in a linear range for glucose detection from 2 to 12 mM, a sensitivity of 15.31 $\mu$A mM$^{-1}$ cm$^{-2}$, and LOD equal to 25 $\mu$M. The device exhibited a fast response of around 1 s, remarkable stability for three months when stored at 4 °C and was successfully evaluated in clinical serum samples. Additionally, PAni-based composites have been applied for DET in third-generation biosensors. Jie Zhu et al. developed a PAni-TiO$_2$NT composite, departing from TiO$_2$NT synthesized by hydrothermal method and subsequently polymerizing PAni by COP (Figure 9C) [81]. A suspension of the PAni-TiO$_2$NT composite, GO$_x$ and Nafion was mixed and cast over a GCE to create the glucose biosensor. The electrochemical characterization revealed the DET of immobilized GO$_x$ produced by electron exchange between FAD and the PAni-TiO$_2$NT/GCE electrode. Overall, the biosensor exhibited a narrower linear range than previous generations, from 10 to 2500 $\mu$M, a sensitivity of 11.4 $\mu$A $\mu$M$^{-1}$, and a low LOD of 0.5 $\mu$M. The current decreased less than 10% after one month, which was attributed to the excellent biocompatibility of the composite. Furthermore, the described representative cases, the Table 1 presents a more comprehensive revision from 2015 to date of enzymatic PBGS.

4.2. PAni-Based Implantable Glucose Biosensors

So far, taking glucose measurements at a given time has been the classical approach for monitoring; however, the needs of diabetics healthcare require a focus on a continuous approach. The materials associated with continuous and implantable devices need to exhibit superior biocompatibility as well as avoid the biofouling of the device. In this sense, PAni has been evaluated as a material for in vivo detection of subcutaneous glucose, Table 1. A biosensor presented by Lu Fang et al. was developed with an arrangement of several layers: Electrodeposited Pt layer, a PAni/GO.$x$ layer, Polyurethane (PU) layer, and Epoxy-PU layer (E-PU) [82], Figure 10. In particular, PAni was electropolymerized over the working electrode, and GO.$x$ was immobilized by crosslinking. PU and E-PU provided stability and biocompatibility as well as a protective layer for the implantable sensor. First, the Pt/PAni- GO.$x$/PU/E-PU biosensor was evaluated in PBS and bovine serum, showing similar analytical parameters of sensitivity (63 nA/mM) and working range (0–20 mM). Then, the device was tested for in vivo implantation in rats connecting the electrodes under the skin and recording the signals produced by dextrose injection. The recorded in vivo values showed good accordance with blood glucose obtained from the vein tail, observing a delay of 8 min attributed mainly to physiological phenomena in the implanted tissue. The biosensor showed an in vivo lifetime of 26 days, with a fluctuation period during the first 12 days, followed by a stable period from day 12 to 26. The authors attributed the extensive life to excessive GO.$x$ immobilized in the PAni NF matrix and the protective PU/E-PU membrane. In a different approach, a flexible, double-sided SPCE was used as the base of a PB doped/PAni/GO.$x$/PU biosensor [83]. In this design, the front was manufactured with a carbon ink doped with PB as the working electrode, and the back was printed with Ag/AgCl ink as the counter electrode. Then, the carbon/PB electrode was modified with PAni and crosslinked GO.$x$, and finally covered with a protective PU layer. The cytotoxicity study revealed the benefits of PU coating for the working electrode composite even though it showed minimal effect for the highly toxic Ag/AgCl electrode. The PB doped/PAni/GO.$x$/PU also showed advances in terms of manufacturing; nevertheless, the
in vivo experiments showed that the information recorded by the implantable sensor needs improvements to correlate with registered blood glucose.

Figure 10. Implantable PANi-based biosensors for continuous glucose monitoring: (a) A view of the CAD design of the Pt/PAni/GOx/PU/E-PU biosensor and the real implantable device, (b) SEM view of the PANi NF immobilization matrix, (c) recorded subcutaneous glucose response (blue) for in vivo test against blood glucose measurement, (d,e) fabrication of PB doped/PAni/GOx/PU biosensor, (f) SEM view of GOx immobilized in PANi matrix, (g) implanted device in rat specimen for in vivo test. Images reproduced from references Lu Fang et al. [82] and Yu Cai et al. [83].
Table 1. Enzymatic PANi-based glucose sensors (PBGS).

| Gen | Material | Immob. Method | Polymerization | Technique | Linear Range | Sensitivity $\mu$A·mM$^{-1}$·cm$^{-2}$ | LOD $\mu$M | Sample | Stability | Ref. |
|-----|----------|----------------|----------------|-----------|--------------|--------------------------------------|---------|--------|-----------|------|
| 1st | PANi/AuNPs | crosslinking | rapid mixing | AMP | up to 16.5 mM | 65.4 | 70 | serum | 22 days | [92] |
| 1st | PANiHF/PAnPs | crosslinking | in situ polymerization | AMP | 0–24 mM | 35 $^\text{**}$ | - | blood | 7 days | [77] |
| 1st | PANi | potentiostatic entrapment | ECP | AMP | 0.01–0.1 M | - | - | - | - |
| 1st | NH$_2$-MWCNT/rGO/PAni/AuNPs | covalent bonding | physical absorption | COP | 1–10 mM | 246 | 64 | blood serum | 30 days | [58] |
| 2nd | PANi/PEO | physical absorption | rapid mixing polymerization | ECP | up to 40 mM | - | - | - | - |
| 2nd | PAni/PFO | physical absorption | free-radical polymerization | ECP | 1–12 mM | 15.31 | 25 | serum | 3 months | [59] |
| 2nd | PAni/VPAni/LuPc$_2$ | crosslinking | interfacial polymerization | AMP | 0.5–50 mM | 2.8$^*$ | 89 | serum | 8 days | [73] |
| 2nd | PAA-VS-PAni/GPL-FePc | crosslinking | interfacial polymerization | AMP | 0.1–100 mM | 14.6 mV/decade | - | - | - | [88] |
| 3rd | PAni/rGO | crosslinking | interfacial polymerization | AMP | 0.1–100 mM | 14.6 mV/decade | - | - | - | [89] |
| 3rd | PANi | EC entrapment | potentiometric | ECP | 0.1–100 mM | 14.6 mV/decade | - | - | - | [89] |
| 3rd | PANi/CoC$_2$O$_4$ | covalent bonding | vapor phase polymerization | AMP | 20–140 $\mu$M | 163.14 | 5.33 | - | - | [84] |
| 3rd | PAni/MWCNT/AuNPs | entrapment in chitosan | vapour phase polymerization | COP | 2-13 mM | 0.099 $^*$ | 94 | - | - | [85] |
| 3rd | PAni/activated carbon/TiO$_2$ | crosslinking | in situ polymerization | COP | 0.06–1.19 mM | 29.17 | 0.21 | serum | - | [57] |
| 3rd | PAni microtubes | electrostatic attraction | in situ polymerization | COP | 4–800 $\mu$M | 35.42 | 0.8 | serum | - | [68] |
| 3rd | PAni/TiO$_2$ NT | entrapment | ECP | 10–2500 $\mu$M | 11.4 $^*$ | 0.5 | - | 2 months | [81] |
| 3rd | PAni/PVP-AuNPs/GOx/Nafion | physical absorption | galvanostatic polymerization | AMP | 0.05–2.25 mM | 9.62 | 10 | serum | 2 weeks | [75] |
| 3rd | PAniNW/rGO | electrostatic interaction | galvanostatic polymerization | AMP | 0.1–8.5 mM | 12.3 | 0.1 | - | 25 days | [87] |
| 3rd | TiO$_2$/PAni | entrapment in chitosan | vapor phase polymerization | AMP | 20–140 $\mu$M | 163.14 | 5.33 | - | - | [84] |
| 3rd | PAni/activated carbon/TiO$_2$ | crosslinking | in situ polymerization | AMP | 0.02–6.0 mM | - | 18 | - | - | [86] |
| 3rd | PAni microtubes | electrostatic attraction | in situ polymerization | AMP | 4–800 $\mu$M | 35.42 | 0.8 | serum | - | [68] |
| 3rd | PAni/TiO$_2$ NT | entrapment | ECP | 10–2500 $\mu$M | 11.4 $^*$ | 0.5 | - | 2 months | [81] |
| 3rd | PAni/PVP-AuNPs/GOx/Nafion | physical absorption | galvanostatic polymerization | AMP | 0.1–8.5 mM | 12.3 | 0.1 | - | 25 days | [87] |
| 3rd | PAniNW/rGO | electrostatic interaction | galvanostatic polymerization | AMP | 0.1–8.5 mM | 12.3 | 0.1 | - | 25 days | [87] |

$^*$ LOD provides an approximation for the limit of detection.
Table 1. Cont.

| Gen | Material | Immob. Method | Polymerization | Technique | Linear Range | Sensitivity $\mu A$ mM$^{-1}$ cm$^{-2}$ | LOD $\mu M$ | Sample | Stability | Ref. |
|-----|----------|---------------|----------------|-----------|--------------|-------------------------------|-----------|--------|-----------|------|
| 1st | PAni-Montmonirollite-PtNPs | entrapment in chitosan | ECP | AMP | 0.01–1.94 mM | 35.56 | 0.1 | blood serum | 40 days | [60] |
|      | PB doped ink/PAni/GOx/PU | physical absorption | COP | AMP | 0–12 mM | 16.66 | - | in vivo | 14 days | [83] |
|      | Graphene/PAni-co-PDPA | physical absorption | interfacial polymerization | ECP | AMP | 1–10 $\mu M$ | 0.51 * | serum | 20 days | [64] |
|      | PAN/PAni/Graphene | entrapment in chitosan | COP | AMP | 0.01–1.97 mM | 29.11 | 2.10 | penicillium | 1 month | [61] |
|      | PAni/SnO$_2$NF | entrapment in chitosan | COP | AMP | 5–100 $\mu M$ | - | 1.8 | urine | 60 days | [56] |
|      | PAni-SDS-F127 | crosslinking | in situ polymerization | AMP | 5–50 mM | 485.787 | 3.202 | - | - | [80] |
|      | PAni/PAA | physical absorption | ECP | AMP | 0–16 mM | 49.3 | 26.5 | - | - | [95] |
|      | PAni/SnO$_2$@3D-rGO | physical absorption | CO/Click chemistry | AMP | 0.56 nM–270 $\mu M$ | - | 0.26 nM | serum | 1 month | [63] |
|      | PAni-g-PEG | crosslinking | graft polymerization | AMP | 0.05–1.0 mM | 47.72 | 20 | - | 20 days | [71] |
|      | PAniNFW/nanodiamond | crosslinking | graft polymerization | AMP | 1–30 mM | 2.03 * | 18 | serum | 30 days | [70] |
|      | PAniNFs/GOx/PU/E-PU | crosslinking | covalent bonding | ECP | AMP | 0–20 mM | 63 ** | - | in vivo | 14 days | [82] |
|      | MWCNT-COOH/PAni | physical absorption | COP | FET | 0.005–500 mM | - | 0.5 | serum | 1 week | [96] |
|      | MWCNT/Au NPs/PAni | physical absorption | in situ polymerization | CV | 2–12 mM | 12.73 | - | - | 2 days | [97] |
|      | 4-amino thiophenol/Au NPs/GOx–HRP/6-mercaptop-o-1-hexanol-11-mercaptoundecanoic acid/Au | covalent bonding | enzymatic polymerization | AMP | 0.0165–10.0 mM | 41.78 | 5.4 | serum | 10 days | [98] |
|      | Graphene/PAni | entrapment in chitosan | ECP | AMP | 0.01–1.48 mM | 22.1 | 2.769 | plasma | - | [72] |
|      | 3D PB/PAni/MWCNT | crosslinking | ECP | AMP | 0.05–4 mM | - | 40 | in vivo | 1 h | [99] |
|      | AuNPs/PAni-Pt NPs/GOx/PVDF-Nafion | entrapment in chitosan | EC | AMP | 0–20 mM | 0.23 * | - | in vivo | 21 days | [76] |
|      | PB/IL-PAni/MWCNT | physical absorption | ECP | AMP | 0.0125–1.75 mM | 94.79 | 1.1 | serum | - | [62] |
|      | PAni/k-carrageenan | physical absorption | COP | DPV | 1–15 mM | 0.86 | - | - | - | [100] |

COP: Chemical oxidative polymerization; ECP: Electrochemical polymerization; EC entrapment: Electrochemical entrapment; AMP: Amperometry; CV: Cyclic voltammetry; DPV: Differential pulse voltammetry; * $\mu A$/mM; ** nA/mM. PAA-VS-PAni/GPL-FePc: polyacrilic acid (PAA)-vinyl substituted polyaniline (VS-PAni)/iron phthalocyanine functionalised graphene nanoplatelets (GPl-FePc). SiO$_2$-LuPc$_2$/PAni-PVIA-CNB: lutetium phthalocyanine incorporated silica nanoparticles (SiO$_2$-LuPc$_2$) grafted with (poly(vinyl alcohol-vinyl acetate) itaconic acid) doped polyaniline conducting nanobeads (PAni-PVIA-CNB).
5. Polyaniline-Based Non-Enzymatic Glucose Sensor

The design of novel nanomaterials for non-enzymatic glucose detection has raised considerable attention in recent years. The primary motivation is to avoid the drawbacks associated with enzymatic glucose operation, e.g., limited temperature conditions, narrow pH range, the need for a biocompatible environment, and operational instability after long-term storage. However, the proposed materials aiming to replace the action of GO must ensure a mechanism to replace the high selectivity of the enzyme; such sensors also need to function at physiological pH conditions and provide a proper cost-to-benefit operation. Moreover, as presented in Section 2.1.4, the non-enzymatic glucose sensing mechanisms are mainly described in alkaline conditions. So far, Pt and AuNPs are among the main reported materials for non-enzymatic detection, although transition metals, metal alloys, and carbon-based nanomaterials present promising advances [101]. Similarly, PANi-based for non-enzymatic glucose sensing has been increasingly reported during the last years. For example, PANi as electrocatalytic material for glucose oxidation was evaluated by Ameen et al. with nanocage-augmented PANI nanowires (NCaPANI NWs) [102]. During CV, the sensor showed a considerable current increment after glucose addition, which was attributed to the ability of the NCaPANI NW surface to adsorb a large number of glucose molecules. While the detection was performed in a PBS buffer at pH 7.0, the working range was limited to 10–120 µM. Table 2 displays other examples of non-enzymatic PBGS with PANi as sole transducer.

For improving the analytical features of detection, PANi combined with metallic nanoparticles (MNPs) has been applied for detection, Table 3. An essential function of MNPs is to increase the electronic transfer capacity of the PANi towards the electrode because this interaction enhances both the intrinsic conductivity and the surface area of the PANi. The electrocatalytic activity of MNPs is another elementary function in the design of glucose sensors. Its use allows solving some of the problems implied by enzymes, such as sensitivity to pH, temperature, the element of detection, and others. In this sense, a hybrid material, PANi@CuNi, was proposed by Bilal et al. [103] and obtained by mixing in the solution PANi synthesized by inverse emulsion polymerization and CuNi NPs synthesized by polyol method (Figure 11). The PANi provides a high surface area for NP growth, providing high catalytic activity towards glucose oxidation due to Ni oxidation states. The sensor exhibited a linear range from 0.1 to 5.6 mM, a sensitivity of 1030 µA mM⁻¹ cm⁻², and LOD of 0.2 µM. While the device was evaluated with blood serum samples, additional conditioning with an alkaline medium was necessary for detection. In a different approach, Yassin et al. [104] selected Co₃O₄, a semiconductor with electrocatalytic behavior and chemical stability, for a novel non-enzymatic sensor. Co₃O₄ showed a nanosheet morphology, subsequently incorporated in PANiNFs. The Co₃O₄@PANiNFs were cast onto the GCE, showing excellent electrical conductivity and a large active area. The addition of PANi allows a fast electron transfer and reversible redox process. An essential feature of this device is that glucose detection was performed in 0.1 M PBS (pH 7.4) as an electrolyte. The authors proposed that the registered anodic current due to glucose oxidation could be related to synergistic effects between PANi states and Co3O4 nanosheets. Additionally, the Co₃O₄@PANiNFs sensor achieved a linear detection range from 0.1 to 8 mM, operational stability after 27 days, and the ability of detection in blood serum samples.

The development of 3D structures increases the surface area and improves the electronic transmission efficiency of the electrode, Table 4. For example, Fang et al. obtained a 3D structure of PANi and rGO (Figure 12) [105]. PANi and GO were co-deposited by ECP on the electrode; then GO was electrochemically reduced to rGO, and the addition of electrodeposited CuO NPs completed the composite. Because the PANi conductivity is strongly influenced by pH, the alkaline electrolyte solution significantly impacts the performance of the composite, as observed by the authors in a composite without rGO. Thus, the combination of PANi with carbon nanomaterials improves the electrocatalytic activity of the proposed sensor, enhances the analytical parameters (linear range 0 to 13 mM), and promotes a fast response (<3 s). In a different approach, a multicomponent nanobead (MCNB)
was developed using a copolymer of poly(aniline-co-anthranilic acid)-grafted-graphene (G-PANI(COOH), branched polyethyleneimine (b-PEI), and ferrocene carboxyaldehyde (Fc-CHO). All the components were integrated into a network structure further decorated with CuNPs. The G-PANI(COOH)-PEI-Fc/Cu NP assembly was applied for oxidation of glucose in 0.1 M PBS, allowing detection at lower potential (+0.37 V). The performance was attributed to the synergistic effects of MCNB assembly, which allowed large amounts of glucose absorption among enhanced characteristics. Glucose detection was performed in DPV, achieving a linear range from 0.5 to 14 mM [106].

Figure 11. Non-enzymatic Co$_3$O$_4$@PaniNFs sensor: (a) FE-SEM view of composite, (b) calibration curve for glucose detection, and (c) selectivity of the device; PAni@CuNi sensor: (d) SEM image of the nanocomposite, glucose detection in 0.1 NaOH using LSV (e) and CV (f); The figure was adapted from references Bilal et al. [103] and Yassin et al. [104].

Figure 12. (a) SEM view of PANi/rGO/CuO NP sensor, (b) CV detection of glucose, (c) proposed sensing mechanism, (d) glucose detection with PANi(COOH)-PEI-Fc/Cu NP sensor using DPV and the obtained calibration curve, (e) selectivity of the device against common interferences, (f) performance against the commercial device. The figures were adapted with permissions from references Fang et al. [105] and Gopalan et al. [106].
Table 2. Non-enzymatic PAni-based glucose biosensors.

| Material Electrode | Polymerization | Technique | Solution-pH | Linear Range | Sensitivity | LOD | Ref. |
|--------------------|----------------|-----------|-------------|--------------|-------------|-----|------|
| CPAniNS 1          | COP            | AMP       | 0.1 M H₂SO₄| 1–1000 μM   | 2003.5 μA·mM⁻¹·cm⁻² | 0.043 μM | [107]|
| PAni/BioHAP/AgHg 2  | COP            | AMP       | 0.1 M PBS-7-4| 1.0–10.0 mM | -           | 1.0 mM | [108]|
| NCa-PAni NWs 3      | ECP            | LSV 4     | 0.1 M PBS-7 | 10–120 μM   | 156.4 μA·mM⁻¹·cm⁻² | 0.657 μM | [102]|

1 Conductive Polyaniline Nanosheets; 2 PAni/biomineralized hydroxyapatite/silver amalgam; 3 nanocages-augmented PAni nanowires; 4 LSV Linear Sweep Voltammetry.

Table 3. Non-enzymatic PAni/nanomaterials based glucose biosensors.

| Material Electrode | Polymerization | Technique | Electrolyte | Linear Range | Sensitivity | LOD | Sample | Ref. |
|--------------------|----------------|-----------|-------------|--------------|-------------|-----|--------|------|
| NiNPs/PAni        | ECP            | AMP       | 0.1 M NaOH  | 0.02–10 mM   | 278.8 μA·mM⁻¹·cm⁻² | 1 μM | -      | [109]|
| CuNPs-Halloysite NT/PAni | COP      | AMP       | 0.1 M NaOH  | 0.01–0.1; 0.1–0.5 mM | 434.0; 57.7 μA·mM⁻¹·cm⁻² | 0.27 μM | -      | [110]|
| PAni/Cd/STN        | COP            | AMP       | 0.1 M NaOH  | 0.0005–0.01 M | 5.23 μA·mM⁻¹·cm⁻² | 1.32 μA·mM⁻¹·cm⁻² | 0.03 mM and 0.5 mM | [111]|
| NiFeNPs/PAni       | COP            | AMP       | 0.1 M NaOH  | 0.02–1 mM    | 1050 μA·mM⁻¹·cm⁻² | 0.5 μM | -      | [112]|
| Polyaacronitrel/    | COP            | AMP       | 0.1 M NaOH  | 100 μM–33 mM | 360 μA·mM⁻¹·cm⁻² | 1.2 μM | -      | [113]|
| PAni/CuO           | COP            | AMP       | 0.1 M NaOH  | 0.01–10 mM   | 379.8 μA·mM⁻¹·cm⁻² | 0.15 μM | -      | [114]|
| AuNPs-TiO₂/PAni    | ECP            | AMP       | 0.1 M NaOH  | 1–3000 μM    | 5625 μA·mM⁻¹·cm⁻² | 0.19 μM | serum | [115]|
| NiO/PAni           | COP            | AMP       | 0.1 M NaOH  | 5–9800 μM    | 146.6 μA·mM⁻¹·cm⁻² | 0.06 μM | serum | [116]|
| PdNW/3D-PAni       | COP            | AMP       | 0.1 M NaOH  | 0.01–8 mM    | 313.6 μA·mM⁻¹·cm⁻² | 0.15 μM | serum | [117]|
| AuNPs-TiO₂/PAni    | ECP            | AMP       | 0.1 M NaOH  | 1–3000 μM    | 5625 μA·mM⁻¹·cm⁻² | 0.19 μM | serum | [118]|
| PAni/Ag nanoleaf   | COP            | AMP       | 0.1 M NaOH  | 1–3000 μM    | 5625 μA·mM⁻¹·cm⁻² | 0.19 μM | serum | [119]|
| Cu-PAni            | ECP            | AMP       | 0.1 M NaOH  | 0.001–1; 1–10 mM | 1625; 1325 μA·mM⁻¹·cm⁻² | 130 nM | -      | [120]|
| Ni-SoO₃/PAni/      | ECP            | AMP       | 0.1 M NaOH  | 1–3000 μM    | 5625 μA·mM⁻¹·cm⁻² | 0.19 μM | serum | [121]|
| CuO/cotton         | COP            | AMP       | 0.1 M NaOH  | 0.001–1; 1–10 mM | 1625; 1325 μA·mM⁻¹·cm⁻² | 130 nM | -      | [121]|
| Co₃O₄@PAniNFs      | COP            | AMP       | 0.1 M LiOH  | 1.0 μM–20 μM | 237 μA·mM⁻¹·cm⁻² | 0.06 μM | serum | [122]|
| PAni/NiF²          | COP            | AMP       | 0.1 M NaOH  | 0.001–19.899 | 1030 μA·mM⁻¹·cm⁻² | 0.2 μM | serum | [108]|
| Hollow CuO/PAni NHz | COP            | AMP       | 0.1 M NaOH  | 0.001–19.899 | 1030 μA·mM⁻¹·cm⁻² | 0.2 μM | serum | [108]|
| PAni/CuNi          | ECP            | AMP       | 0.1 M NaOH  | 0.02–10 mM   | 278.8 μA·mM⁻¹·cm⁻² | 1 μM | -      | [109]|
| CuNPs/PAni         | COP            | AMP       | 0.1 M NaOH  | 0.02–10 mM   | 278.8 μA·mM⁻¹·cm⁻² | 1 μM | -      | [109]|
| NiO/CuO/PAni       | ECP            | AMP       | 0.1 M NaOH  | 20–2500 μM   | 2 μM        | 2 μM | serum | [125]|
| NiCO3@PAni         | COP            | AMP       | 0.1 M NaOH  | 0.01–4.7350 mM | 4.55 μA·mM⁻¹·cm⁻² | 0.383 μM | -      | [126]|
| PAni/AuNPs         | -              | EIS       | K₃[Fe(CN)₆]₃/0.1 M PBS | 0.3–10 mM | - | 0.1 mM | - | [127]|

1 PAni nanosheets; 2 Nickel foam; 3 PAni nanohybrid fibers.
### Table 4. Non-enzymatic PAni/Carbon-based materials/nanomaterials glucose biosensors.

| Material Electrode                  | Electrode | Polymerization | Technique  | Electrolyte | Linear Range | Sensitivity       | LOD        | Sample         | Ref.  |
|------------------------------------|-----------|----------------|------------|-------------|---------------|-----------------|------------|----------------|-------|
| ZnCoO$_4$/g-C$_3$N$_4$/Pani$^1$    | COP       | AMP            | -          | 0.1–3 mM    | 15.64 mA-mM$^{-1}$·cm$^{-2}$ | 4 µM       | -              | [128] |
| PLA/GR/PAni/Cu                     | ECP       | AMP            | 0.1 M KOH  | 1–7 mmol-L$^{-1}$ | -             | 49.3 mmol-L$^{-1}$- | -          | [129] |
| CuNWs/PAni/rGO                    | -         | -              | -          | 0–4 mM      | 843.06 µA-mM$^{-1}$·cm$^{-2}$ | 1.6 mM     | -              | [130] |
| Cu$^{2+}$/PAni/rGO ink             | COP       | AMP            | 0.1 M NaOH | 2.8–22.2 µM; 0–4 mM | 4168.37; 525.4 µA-mM$^{-1}$·cm$^{-2}$ | 4.93 µM serum | [131] |
| PAniNS@rGO                        | COP       | AMP            | 0.1 M NaOH | 1–10 mM     | 3448.27 mA-mM$^{-1}$·cm$^{-2}$ | 30 nM      | -              | [132] |
| TiO$_2$rGO-PAni                   | COP       | LSV            | 0.1 M NaOH | 10–180 µM   | -             | 7.46 µM         | -          | [133] |
| PAni/rGO/CuO                      | ECP       | AMP            | 0.1 M NaOH | 0–13 mM     | 1252 µA-mM$^{-1}$·cm$^{-2}$ | 1.5 µM blood serum | [105] |
| PAni/ZnO/MWCNT                    | ECP       | LSV            | 10 mM NaOH | 0.1–1; 1–6 mM | 7.8307; 1.6731 µA-mM$^{-1}$·cm$^{-2}$ | 0.1 µM - | [134] |
| fGO/Fe$_2$O$_3$/PAni$^2$          | COP       | CV             | PBS        | 0.05 µM–5 mM | -             | 0.01 µM serum  | -          | [133] |
| NiCo$_2$S$_4$/rGO@PAni            | COP       | AMP            | 0.1 M NaOH | 1–7000 µM   | 4.924 µA-mM$^{-1}$·cm$^{-2}$ | 0.14 µM serum | -          | [135] |
| NiO/Au/PAni/rGO                   | ECP       | AMP            | 0.1 M NaOH | 0.09–6 mM   | -             | 0.23 µM serum   | -          | [136] |
| AuNPs/PAni                        | ECP       | CV             | 0.5 M KOH  | 10.26 µM–10.0 mM | 150 µA-cm$^{-2}$·mM$^{-1}$ | 3.08 µM | serum          | [137] |
| PAni/Co$_2$O$_3$ nanorods         | -         | AMP            | 0.1 M KCl  | Up to 0.6 mM | 25.79 µA-mM$^{-1}$·cm$^{-2}$ | 0.56 µM - | [138] |
| CuNPs/PAni/graphene               | COP       | AMP            | 0.1 M NaOH | 0.001–3.7 mM | 150 µA-cm$^{-2}$·M$^{-1}$ | 0.27 µM mice serum | [139] |
| G-PAni(COOH)-PEI-Fc/Cu-MCNB       | COP       | DPV            | 0.01 M PBS | 0.50–14 mM  | 14.3 µA-mM$^{-1}$·cm$^{-2}$ | 0.16 mM serum | [130] |
| NiONPs/PAniNW/GO                  | ECP       | AMP            | 0.1 M NaOH | 2 µM–5.560 mM | 376.22 µA-mM$^{-1}$·cm$^{-2}$ | 0.5 µM bovine serum | [140] |

$^1$ Zinc cobaltite/graphitic carbon nitride/PAni; $^2$ carboxy functionalized graphene oxide (fGO).
6. Conclusions and Future Directions

Glucose monitoring is an essential tool for the daily care of DM patients, and thus the development of accurate devices for detection is of high interest. PANi is a versatile CP with favorable characteristics for biomedical applications. So far, the majority of reviewed literature related to PANi applications in DM is focused on glucose sensing, using either enzymatic or non-enzymatic mechanisms. This is attributable to the intrinsic characteristics of PANi such as conductivity, biocompatibility, and tunable redox behavior, which allow fast, sensitive, and stable sensing devices. However, some significant challenges need to be addressed to propel PANi-based devices into a reality in DM care. Firstly, most of the sensors have been proposed from a material science perspective, rather than having a specific design to fit the parameters of biological samples. For example, blood glucose concentration in healthy individuals has been set in 5.6 mM, but for self-monitoring of diabetic patients, values up to 7 mM can be expected. While most of the reported PBGS presented an adequate LOD, the majority needed to increase their working range to fit the needs of blood glucose analysis. Specifically for non-enzymatic sensors, a key challenge is the dependence of alkaline electrolytes for glucose detection, which adds additional pre-processing steps and significantly impacts PANi conductivity. In this sense, non-enzymatic PBGS for detection in PBS similar to physiological conditions have been presented and need to be further improved. A growing and desirable field for DM is the continuous monitoring of glucose. PBGS have been developed for this purpose, achieving promising results for in vivo experiments. However, additional work for improving the biocompatibility of the entire electrochemical device needs to be addressed. A subject for further research is the trend of non-invasive, wearable devices capable of quantifying glucose in different body fluids such as urine, sweat, tears, and saliva. These alternative biological samples pose additional challenges such as low glucose concentration, a more comprehensive range of pH conditions, limited sample collection, and additional interferences. Few works related to the thematic have been presented for PBGS, but given the advances in manufacturing techniques, the polymer could be incorporated in novel wearable devices for on-body glucose detection. Besides glucose monitoring, the ability of PANi for switching between doped and undoped forms for acetone detection, one of the three ketone bodies related to diabetic ketoacidosis, has been explored. This property could be exploited for miniaturized sensors for both analytes with PANi as base material. Overall, this review has presented an overview of the current status of PBGS and the future directions for propelling this technology into DM care devices.

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