

Biosensor strategies to detect serum glycobiomarkers**ABSTRACT**

Changes in glycosylation profiles have emerged as indicators of diseases. Altered glycans and glycoproteins secreted by pathological tissues are found in human serum and are potential glycobiomarkers for early diagnostic and prognostic of diseases such as inflammation, infection and cancer. To obtain serum samples is minimally invasive and a simple procedure; then detection of glycobiomarkers in serum is attractive for clinical applications. Biosensors are a friendly strategy for rapid, easy and highly sensitive measuring of glycoprotein biomarkers. The use of lectins as biorecognition elements in glycobiosensors has provided a specific detection and profiling of glycans linked to glycoproteins. Electrochemical glycobiosensors based in lectin interactions have been employed to characterize glycan profiles in serum glycoproteins and constitute a promising tool in diagnostic and monitoring of diseases.

Keywords: glycosylation; glycans; biomarkers; serum glycoproteins; biosensors; lectin biosensors.

1. INTRODUCTION

Glycosylation is a post-translational modification occurring in more than half of human proteome and glycans have important roles in physiological functions and diseases. Changes in glycosylation profiles are frequently observed on cell surfaces and serum glycoproteins in the occurrence of pathological processes, such as immune disorders, infectious diseases and cancer. Evaluations by chromatographic techniques, mass spectrometry, among other approaches in tissues, cells, biological fluids and serum samples reveal quantitative and structural alterations in glycans in diseases when compared with healthy samples [1, 2]. Then, correlation of some glycoproteins and glycans with diseases make them glycobiomarkers of choice for clinical diagnosis and monitoring of prognosis [3].

The diagnostic of diseases and monitoring during treatment through detection of glycobiomarkers present in bloodstream are attractive in terms of low invasiveness, agility and easily. Analytical techniques based in Enzyme Linked Immunosorbent Assay (ELISA) and other immunoassays are most used to detect and measure serological glycobiomarkers in clinical laboratories. In addition, biosensing technology has been applied successfully to identify and quantify glycobiomarkers with high specificity and selectivity, based on biorecognition events connected to a transduction system [4]. Biosensors can be constructed using a variety of transducer types, including optical, piezoelectric and electrochemical, to generate the results. Electrochemical biosensors have been detached since they provide rapid diagnostic by a simple manipulation and low cost, bypassing some limitations showed in other technologies.

Lectins, a group of sugar-binding proteins have been largely employed for biorecognition of sugars in binding assays. Lectins recognize carbohydrates with specificity for monosaccharides or oligosaccharides, free or bound to a glycosylation site, and may distinguish the linkage type. The lectin-carbohydrate binding provides quantitative and qualitative information about expression of glycans in biological samples, being valuable tools to characterize and identify changes in secreted glycoproteins [5]. Techniques based on chromatography, electrophoresis, immunoassays and biosensors use lectins as biorecognition agents. Lectin biosensors contain a sensing platform with immobilized lectins to detect glycobiomarkers in serum samples through lectin-glycan interaction with high specificity [6]. Various groups of research have

46 developed lectin biosensors for cancer, infectious diseases and virus or bacterial glycan detections. Lectins
47 of different sources and specificity groups are employed in biosensing due to their versatility and diversity.

48 Isoforms from a seed lectin of *Cratylia mollis*, denominated Cramoll, belongs to the specificity group
49 of mannose/glucose such as Concanavalin A, Con A [7]. Cramoll biosensors have demonstrated potential to
50 recognize profiles of glycoproteins of different dengue serotypes in human serum [8, 9] and glycans of
51 bacterial lipopolysaccharides [10], through electrochemical methods, representing a good alternative for
52 detection of diseases. This review introduces some advances in the lectin biosensor area directed for
53 detection of glycomarkers to help in the early diagnostic and monitoring disease assays.

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55 2. GLYCOSYLATION

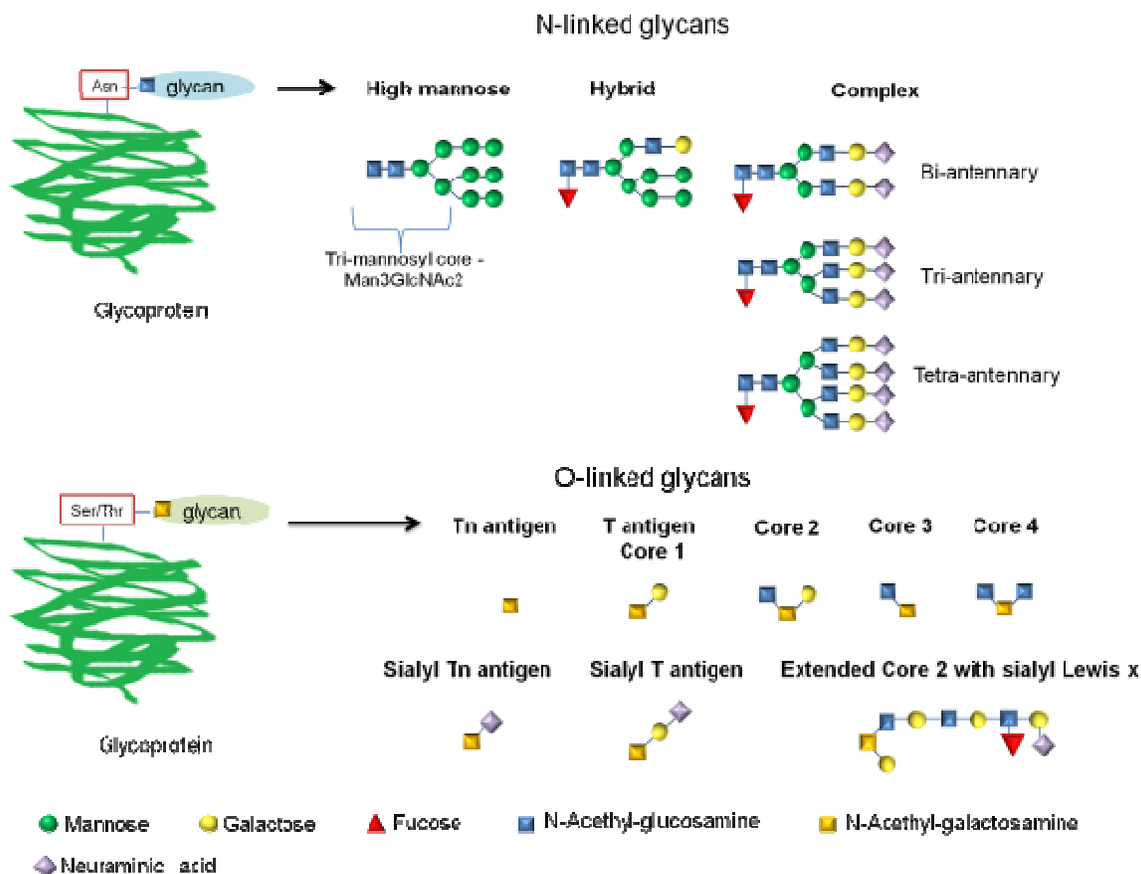
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57 Glycosylation is the most common post-translational modification of proteins and has fundamental
58 importance in biological processes in eukaryotic organisms. It is predicted that nearly 80% of the human
59 plasma proteins are glycosylated [11], in addition to glycosylated proteins secreted and glycan covered cell
60 membranes. Glycosylation reactions involve the covalent attachment of glycan chains to specific amino acid
61 moieties of proteins during and after translation of polypeptide chains in order to form glycoproteins. The
62 glycosylation profile of a protein is determined by a wide group of enzymes known as glycosidase and
63 transferase, residents in the endoplasmic reticulum and Golgi apparatus, which catalyze the extension and
64 transfer of glycan chains to glycosylation site of the protein [12]. They constitute a specific group to a cell
65 type, tissue and organism, resulting in tissue- or cell-specific differences in glycosylation between sites of the
66 same protein [13].

67 Glycans are linked to human glycoproteins *via* two pathways, N- and O-glycosylation, with various
68 branching points. O-linked glycans are attached through hydroxyl group of serine or threonine residues by
69 starting with the addition of N-acetyl-galactosamine (GalNAc –O– Ser/Thr) transferred by an N-acetyl-
70 galactosaminyltransferase in the Golgi apparatus [14]. After, specific transferases elongate different types of
71 core structures, including mucin-type O-linked glycans (core 1, 2, 3, 4; as well as T, TF and Tn antigens), O-
72 linked GlcNAc and O-linked Fuc (Figure 1). N-linked glycans are attached to the amidic nitrogen of
73 asparagine residues within the Asn-X-Ser/Thr glycosylation site, being X different of the proline [15]. N-
74 glycosylation is initiated in endoplasmic reticulum by addition of the precursor tri-mannosyl core
75 Man3GlcNAc2 that is prolonged to generate three subtypes of N-glycans: high-mannose, complex and
76 hybrid (Figure 1). The tri-mannosyl portion of precursor is cleaved by glycosidases and transferred to the
77 Golgi apparatus, where other monosaccharides are added by specific transferases to prolong the glycan
78 chains, generating complex branching. The resulting structures provide the subtypes of complex N-glycans,
79 which have complex branching without mannose residues and hybrid N-glycans, that contain mannose
80 residues and complex branching [15, 16]. In addition, the subtype high-mannose N-glycans contain only
81 mannose residues linked to the tri-mannosyl core [16]. The monosaccharide residues mannose (Man),
82 glucose (Glc), galactose (Gal), fucose (Fuc), N-acethyl-galactosamine (GalNAc), N-acethyl-glucosamine
83 (GlcNAc) and sialic acid (SA) or neuraminic acid (NeuNAc) are the most frequent in the N- and O-glycan
84 chains attached in human proteins [15, 16].

85 Inherent heterogeneity and diversity of glycan structures controlled by enzymes contribute to the
86 varied functions that glycosylation carry out in cells and tissues. Glycosylation plays relevant influence in

87 protein synthesis, processing and function, including folding, stability of tertiary structure, protection against
 88 action of proteases, increased serum half-life of proteins and reduced nonspecific protein-protein interactions
 89 [17, 18, 19]. Moreover, glycosylation mediate a regulation role in many biological processes involving cell-
 90 cell and cell-matrix interactions, such as cell proliferation, cell recognition, adhesion, host-pathogen
 91 recognition, receptor binding, signaling, fertilization, inflammation and immune responses [19, 20].
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Figure 1. Types of N- and O- linked glycans found in human serum.

97 3. GLYCANS ARE POTENTIAL GLYCOBIOMARKERS IN DISEASES

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 99 The activities of glycosylation enzymes are factors dependent modulated by cellular dynamics.
 100 Changes in cellular environment and physiology often due to diseases, affect enzyme activities, resulting in
 101 aberrant glycosylation of proteins [21]. Aberrant glycosylation has been linked to various pathologies such as
 102 infection, chronic inflammations, immunological and genetic disturbs, cancer and metastasis [12, 21, 22].
 103 This involves changing of glycosylation sites, increase or decrease of the site numbers, modification in the
 104 chemical composition or type of glycan, extra branching of glycans and quantitative alteration of specific
 105 glycoprotein on the cell surface and secreted in bloodstream [23, 24].

106 There is strong evidence that these glycosylation changes in the serum glycome are a good way to
107 identify potential glycomarkers of diagnostic and progression of pathological states. Serum glycoproteins
108 such as immunoglobulins, fetuin, haptoglobin, transferrin, alpha-fetoprotein, and other acute phase proteins
109 showed glycosylation changes in response to inflammatory and immune diseases. Immunoglobulin G
110 contains complex bi-antennary N-glycans which have a significant decrease in galactosylation for patients
111 with inflammatory arthritis [25, 26]. The serum glycan profiling of fibrosis patients with hepatitis B and C virus
112 infection revealed a significant decrease of bi-antennary and tri-antennary N-glycans as glycomarkers to
113 monitor the progress of fibrosis [27]. Multi-branched glycans highly sialylated were detected in elevated
114 levels in serum of ulcerative colitis patients correlated with disease degree [28]. Alpha 1-6-linked arm
115 monogalactosylated and core fucosylated bi-antennary N-glycans were reduced in serum glycoproteins from
116 type 2 diabetes mellitus subjects and provide a serological alternative glycomarker [29]. Glycan profiles of
117 whole serum of patients with autoimmune pancreatitis showed elevated levels of agalactosyl and
118 monogalactosyl bi-antennary glycans, being potential biomarkers of the disease [30].

119 Studies have evaluated the presence of aberrant glycoforms and quantitative levels of glycosylation
120 in patients with cancer and compared with benign disease and healthy individuals to identify cancer
121 glycomarkers. Glycosylation aberrant patterns of proteins from cancer cells, tissues and serum commonly
122 involve altered sialylation and fucosylation, change in glycan size and branching, Lewis antigens and
123 truncated O-glycans [12]. Lewis antigens (Figure 2) are frequently expressed on membrane glycoproteins of
124 several cancer cells. Cholangiocarcinoma cells expressed highly sialyl Lewis a ($\text{Neu5Ac}\alpha\text{2-3Gal}\beta\text{1-3(Fuc}\alpha\text{1-4)GlcNAc}$) and glycoprotein mucin 5AC was elevated in tissues and serum from cholangiocarcinoma patients
125 [31, 32]. Serological N-glycome in breast cancer patients showed increased levels of sialyl Lewis x
126 ($\text{Neu5Ac}\alpha\text{2-3Gal}\beta\text{1-4(Fuc}\alpha\text{1-3)GlcNAc}$) antigen and an increased sialylation [33]. Increase of $\beta\text{1-6GlcNAc}$
127 branches in N-glycans added by N-acetylglucosaminyltransferase V (GnT-V) is showed in breast and ovarian
128 cancer [23, 34]. Glycans of prostate-specific antigen isolated from tumoral cell and sera showed tri- and
129 tetra-antennary structures of N-glycans [35]. In cancer cell-surface, O-linked glycans of mucin-type
130 glycoproteins are often defective, resulting in the expression of truncated O-linked glycans denominated of T
131 ($\text{Gal}\beta\text{1-3-GalNAc}\alpha\text{-O-Ser/Thr}$) and Tn ($\text{GalNAc}\alpha\text{1-O-Ser/Thr}$) antigens (Figure 1), as observed in breast and
132 colon cancer [23, 36]. T/Tn antigen quantitative assays were developed to detect cancer without previous
133 biopsy using highly purified T antigen for determination of anti-T immunoglobulins in serum samples [37].
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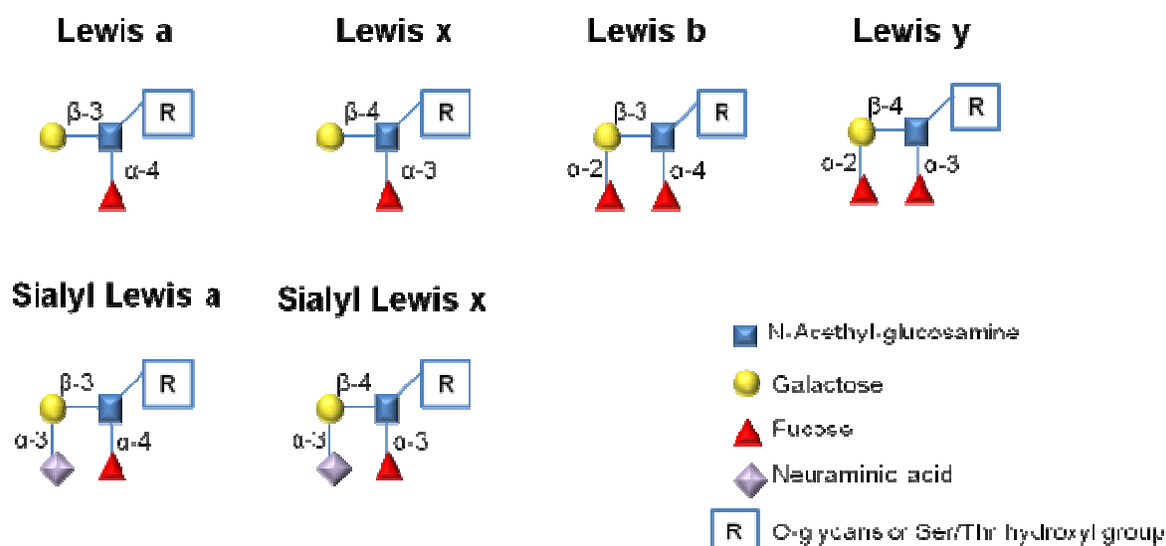


Figure 2. Terminal Lewis and sialylated Lewis antigens expressed by human tumors.

4. SERUM GLYCOPROTEINS AS GLYCOBIOMARKERS OF DISEASES

Clinical tests using body fluids such as serum, plasma, urine and saliva are minimally invasive, easily handled, with short time response, then they are preferred for detection and prognostic evaluation of various diseases. Glycoproteins secreted in the bloodstream are a considerable part of the serum biomarkers and they might show differential quantitative and qualitative level between healthy and pathological samples [3, 38]. Some glycoproteins are well known serologic glyco-biomarkers for cancer diagnostic that may be detected by laboratory clinical assays, such as the carbohydrate antigens (CA) CA-125 for ovarian cancer, CA15-3 for breast cancer, carcinoembryonic antigen (CEA) for colorectal cancer, alpha-fetoprotein (AFP) for hepatocellular carcinoma (HCC) and prostate-specific antigen (PSA) for prostate cancer (Figure 3). The early diagnosis of infectious diseases based in serum glycoproteins is also possible. Dengue virus nonstructural 1 (NS1) antigen is a test commercially available for dengue infection. NS1 antigen is a glycoprotein present in elevated levels in the serum of infected individuals before specific antibodies emerge [39].

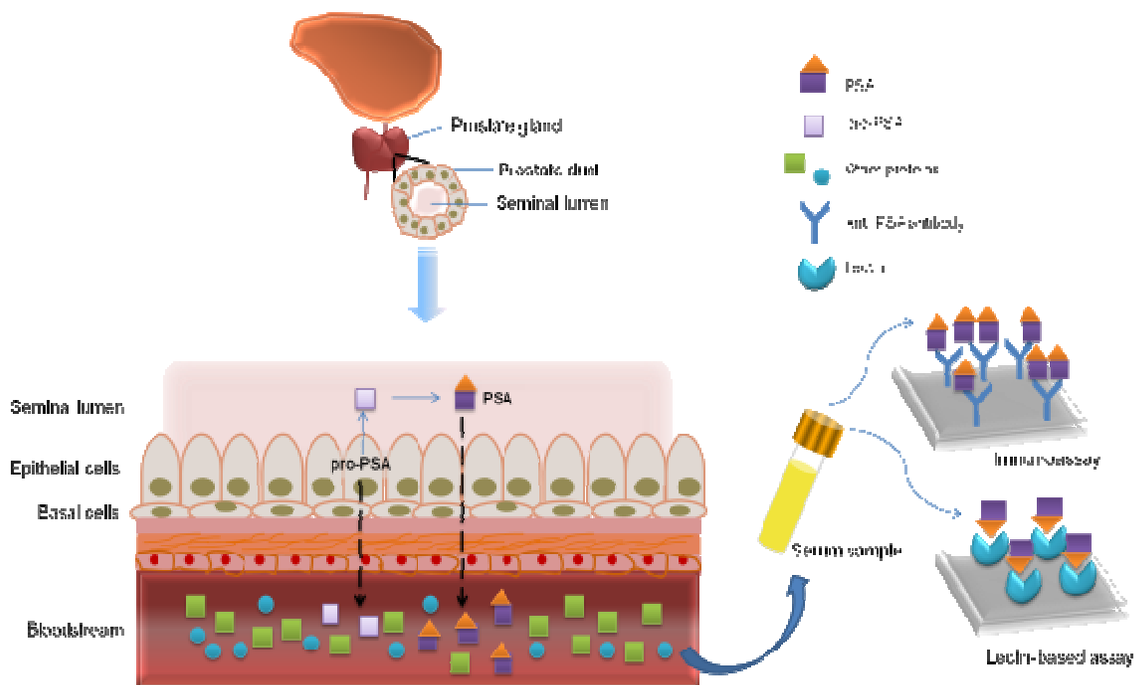
Current tendencies evaluate changes in the glycosylation patterns of serologic glyco-biomarkers appointing them as potential candidates for highly specific biomarkers, for early detection and staging of various types of cancers. Researches appoint an increase of core-fucosylation levels of serum AFP of patients with HCC as improvement of specificity for HCC staging [40]. Altered profiles of fucosylation and sialylation in PSA glycans are reported for prostate cancer and fucosylated PSA showed to have potential to substitute the PSA test in the differentiation of aggressive tumors from non-aggressive tumors of prostate [41]. Neoglycoforms of CEA in cancer identified as bi-antennary, tri-antennary and tetra-antennary glycans with residues of sialic acid and fucose can improve the tumor diagnosis and staging [42]. Haptoglobin is an acute phase glycoprotein produced by the liver and its concentration is elevated significantly in many diseases, including hepatic inflammation, hepatitis and various types of cancer [43, 44]. Studies on altered glycosylation of serum haptoglobin observed high levels of fucosylation in haptoglobin associated with cancer and fucosylated haptoglobin (Fuc-Hpt) is reported as a biomarker for diagnostic and prognosis of pancreatic cancer and colorectal cancer [45]. Elevated levels of serum Fuc-Hpt also have been associated

166 with prostate cancer. An increase of glycan fucosylated bi-, tri- and tetra-antennaries of serum haptoglobin
 167 was observed in prostate cancer patients, correlated with Gleason score [46].

168 Analytical techniques have allowed the search and characterization of neoglycoforms present in the
 169 bloodstream. The structural characterization of glycans involves analyses by robust techniques, such as
 170 liquid chromatography and mass spectrometry, which provide detailed information about composition and
 171 quantity of residues [47]. However, they require long time of analyses, previous treatment of samples,
 172 technical prepare for manipulation, are complex and expensive; then, they become not viable for direct
 173 diagnostic of diseases and monitoring of patients in the clinical treatment. Besides, human serum and
 174 plasma are considered complex samples, because they contain a mixture of proteins, some very abundant
 175 (more than mg/mL) such as albumin, that can mask other protein expressed in low concentrations (ng/mL
 176 and pg/mL range), mainly biomarkers that appear in early diseases.

177 Techniques based in specific binding assays using biorecognition agents (eg. antibody, lectin)
 178 recognize a glycoprotein biomarker or glycan chains linked to protein, measuring these molecules at very
 179 low concentrations in human serum and plasma (Figure 3) with specificity and selectivity [45, 48].
 180 Immunoassays as ELISA commercially available and lectin-based assays have been used for detection and
 181 measurement of specific serum glycobiomarkers in research and clinical applications to diagnostic of
 182 diseases [48, 49]. These techniques frequently require significant sample volumes and several steps for
 183 obtaining results. Biosensors based in binding assays have emerged as a rapid, simple and sensitive
 184 strategy for serological glycobiomarker measurements [50, 51].

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188 **Figure 3. Esquematic representation of secretion from specific glycobiomarker of prostate, the**
 189 **prostate-specific antigen (PSA), in the bloodstream and assays commonly used for specific**
 190 **detection.**

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5. BIOSENSORS: A STRATEGY FOR DETECTION OF GLYCOBIOMARKERS

Biosensors are analytical devices capable of detecting and measuring a specific analyte using a biological recognition element contacted with a transducer, that converts the recognition event into a signal analytically useful and measurable [52]. A biosensor contains a biorecognition element (bioreceptor) that binds specifically to analyte of interest; a transducer element that detect signals (current, voltage, changes of mass, light or temperature) resultant of the interaction bioreceptor-analyte and convert in an electric signal. Besides, the transducer is connected with an appropriate interface for processing of electric signal and display of measurable results [53] (Figure 4).

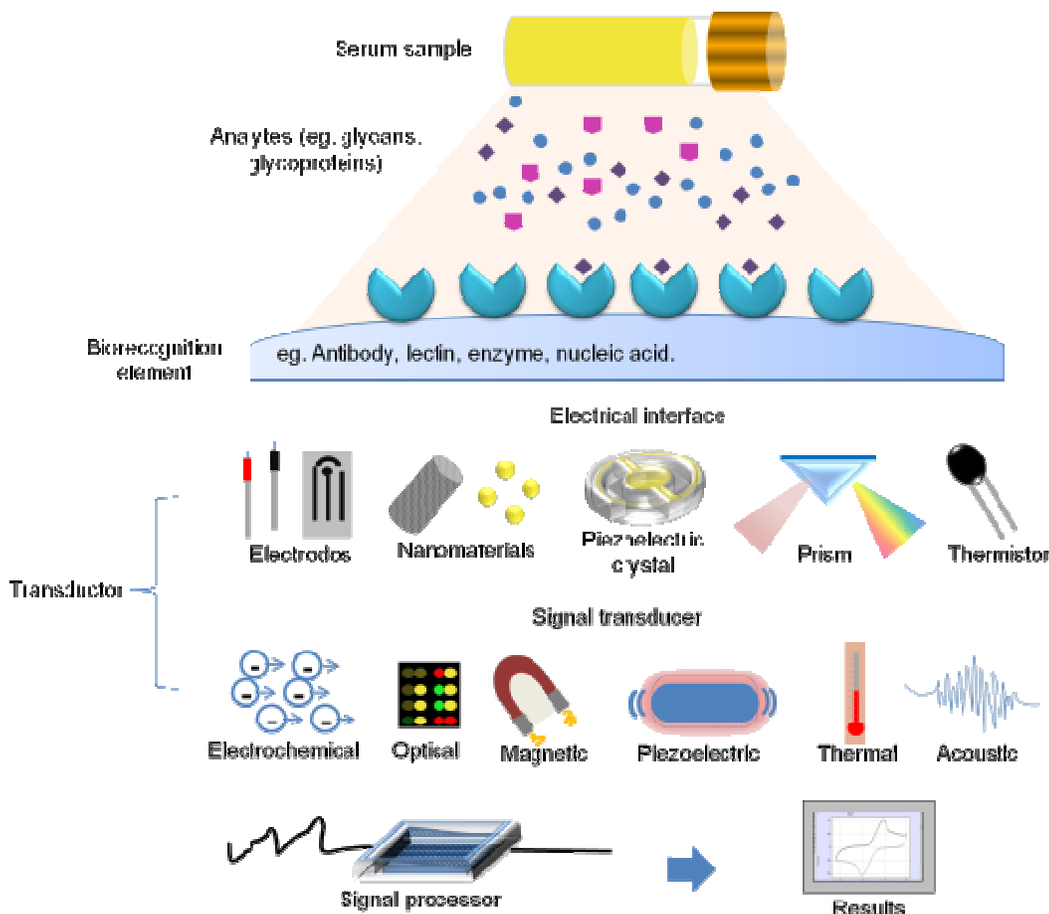


Figure 4. Scheme of a biosensor with a biorecognition element immobilized on the solid surface in direct contact with an electrical interface of the transducer and a signal processor.

According to type of interaction bioreceptor-analyte, the analytical devices are classified as catalytic or affinity biosensors. Catalytic biosensors are based in enzyme activity with generation of product [54]. Affinity biosensors involve binding event between the bioreceptor and analyte without chemical transformations, such as antigen-antibody, nucleic acid hybridization and interaction lectin-carbohydrates [53, 55]. Biosensors are also classified in relation to the signal detected by transducer (Figure 4). They can include an electrochemical biosensor, which detect current, voltage, potential or impedance; an optical

213 biosensor, which measure the intensity of the light emitted; a piezoelectric or acoustic biosensor, that detect
214 variation of mass, pressure or elasticity, and a thermal biosensor, which detect changes in temperature [53].
215 This variety of mechanisms favors a range of designs and applications of biosensors. Electrochemical
216 biosensors are more attractive and predominant in bioanalyses, due to diversity of electrical interfaces and
217 techniques for measurement that collaborate for high selectivity and sensibility; these sensors are easily of
218 construction, portability, possible miniaturization and low cost [56].

219 The first biosensor was developed by Clark and Lyons in 1962, for detection of glucose by glucose
220 oxidase. They elaborated a system known as enzyme electrode, using the enzyme glucose oxidase
221 immobilized on the polyacrylamide film, onto a platinum electrode surface, and the determination of the
222 glucose concentration was based in measurement of oxygen consumed [57]. After, the Yellow Spring
223 Instrument Company developed the Clark and Lyons's technology to introduce in the market the first
224 biosensor of glucose in 1975 [58]. Since then, the biosensor technology developed of different approaches in
225 the detection mode and elaboration of more efficient devices, aiming the various types of analyses [58].
226 Biosensor technologies focus in the fabrication of self-contained, portable and miniaturized devices and at
227 the same time, in conservation of the good analytical properties, such as speed, high sensitivity and
228 selectivity. Biosensors are then friendly for applications to diverse areas: environmental monitoring, biosafety
229 and assays for clinical applications [59].

230 Biosensors or glycobiosensors have been used to detect glycans and glycoproteins, in analyses of
231 human serum [55]. Different of the complex approaches such as glycoproteomic techniques, that require
232 sophisticated instrumentation and significant volume of sample, biosensor advances resulting in easy
233 detection and measuring of glycans or glycoproteins in minimal volume of sample, simple manipulation,
234 short-time response, high sensitivity and specificity, and friendly to use outside of a laboratory [59, 60].
235 These possibilities are attractive for development of point-of-care tests useful in early diagnostic, staging and
236 monitoring diseases such as cancer. The detection of glycobiomarkers using glycobiosensors frequently use
237 antibodies and/or lectins. A variety of available glycobiosensors uses antibodies directed to protein as
238 biorecognition element for specific detection and measurement of serum glycoprotein biomarkers [56]. Lectin
239 glycobiosensors have been developed for characterization of the glycosylation patterns of glycans on serum
240 glycoproteins correlated to diseases, such as degree of sialylation and fucosylation [60, 61].

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242 **6. LECTINS AS BIORECOGNITION ELEMENTS**

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244 The term lectin is derived from the latin word "lectus" (chosen, selected) [62], to define a group of
245 proteins that showed selectivity in the interaction with carbohydrates. Lectins are sugar-binding proteins of
246 non-immune origin that exhibit specific recognition and reversible bind to free carbohydrates and link to
247 glycoconjugates (eg. glycoprotein and glycolipid). They are involved in crucial physiological events of
248 protein-carbohydrate interactions, such as adhesion and cell migration [63, 64]. Glycobiosensors frequently
249 use lectins as biorecognition elements.

250 The association constant between lectin and monosaccharides ranges from 10^3 to $(5 \times 10^4) \text{ M}^{-1}$ and
251 between lectins and oligosaccharides, 10^4 to 10^7 M^{-1} , included values in the same range found for antibody-
252 antigen bindings and enzyme-substrates [65, 66]. Lectins bind to a specific monosaccharide or
253 oligosaccharide through their binding sites through hydrogen bonds, Van der Waals interactions, and
254 hydrophobic interactions [66, 67]. According to the specificity, lectins are classified in five main groups:

255 mannose and/or glucose, N-acetyl-glucosamine, N-acetyl-galactosamine/galactose, L-fucose and sialic acids
256 [68].

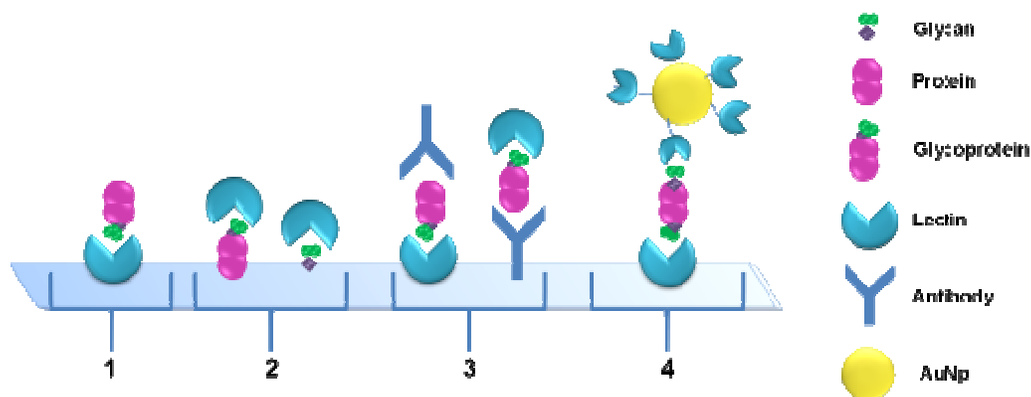
257 Lectins are distributed among viruses, microorganisms and animals although they were initially found
258 in plants [69, 70, 71]. The majority of available lectins are obtained from leguminous plants, comprising the
259 largest number of already characterized proteins, such as Con A, soybean agglutinin (SBA) and *Cratylia*
260 *mollis* lectin (Cramoll). Leguminous lectins, although derived from taxonomically distinct species, have
261 molecular characteristics and physicochemical properties in common, such as Con A obtained from
262 *Canavalia ensiformis* and Cramoll from seeds of *C. mollis*, and they exhibit a great variety in specificity for
263 carbohydrates [67, 72].

264 Lectins have been used in glycoanalytical areas, including analysis of glycosylation profiles and
265 detection of glycobiomarkers in serological samples based in lectin-carbohydrate interaction. They are widely
266 used in the characterization of glycans linked to glycoproteins for comparative analyses between healthy and
267 pathological conditions and identification of aberrant glycan profiles as potential glycobiomarkers [24, 45].
268 According to their specificity, lectins can detect alterations in degree of sialylation [73], fucosylation [46],
269 galactosylation [74], mannosylation [75] and the presence of cancer-associated antigens, like sialyl-Tn (STn)
270 antigen and Lewis antigens [4, 76], and other changes associated with diseases.

271 Glycobiosensors have various lectin-based models to configure the lectin-glycan interactions,
272 sometimes assisted by antibody and nanomaterials to improve the selectivity and signal amplification. The
273 lectin can be directly immobilized onto a solid support to detect glycoproteins by glycan portion, in according
274 to specificity. Following this model, a lectin-based glycobiosensor for profiling of the STn antigen in serum
275 samples was developed by immobilizing the lectin *Sambucus nigra* agglutinin type I (SNA-I) on electrode
276 surface and showed potential to discriminate between cancer and healthy conditions [4]. The inverted
277 configuration based in the immobilization of the glycan or glycoprotein followed by lectin binding also is
278 available. A biosensor based in D-glucose was successful used to evaluate Con A with a limit detection of
279 1.0 pM [77].

280 The two-step sandwich model involves a lectin and an antibody, both to interact with the glycoprotein
281 biomarker. In this case, the capture antibody or lectin is immobilized to recognize the glycoprotein, and in the
282 second step, a lectin or secondary antibody is added to complete the sandwich. A plastic sensor chip
283 based in the technique surface-plasmon field-enhanced fluorescence spectrometry was used to detect
284 GalNAc β 1-4GlcNAc-linked prostate-specific antigen using PSA IgG antibody to capture PSA and the lectin
285 *Wisteria floribunda* agglutinin (WFA) to profile the glycan in serum from patients with benign prostate
286 hyperplasia and patients of prostate cancer [74]. The PSA IgG antibody allowed the characterization of
287 glycosylation profile from the specific glycoprotein target present in serum samples. Other sandwich system
288 is based in the lectin or antibody confined on surface to recognize the glycobiomarker, followed by the
289 addition of lectins coated nanomaterials. The latter compounds are attractive since they increase the amount
290 of secondary biorecognition element and promote the signal amplification. A highly sensitive and selective
291 lectin-based glycobiosensor was developed for comparative analysis of mannose and sialic acid levels on
292 normal and cancer cells using a sandwich system formed by lectins immobilized to detect glycans on cell
293 surface, followed by addition of lectins linked to gold nanoparticles (AuNp) [78]. These strategies provide a
294 characterization of glycosylation profiles on glycoprotein biomarkers with high sensitivity and selectivity,
295 which is useful in clinical diagnostics, including early detection and staging of cancer and monitoring of
296 patients during treatment.

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Figure 5. Configurations of lectin-based assay in glycobiosensing. Direct detection by immobilized lectin (1), reverse detection by incubated lectin (2), lectin-antibody sandwich assay (3) and lectin-conjugate sandwich assay using gold nanoparticles (AuNp) (4).

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307 7 ELECTROCHEMICAL GLYCOBIOSENSORS

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309 Electrochemical methods are the most attractive for sensing biorecognition events due to simple
310 instrumentation, friendly analytical performance and diversity of methods and electrodes available for
311 specific applications [79]. Biosensors based in electrochemical detection are widely applied to glycoanalysis
312 in the field of glycomics to analyze cleaved glycans, together to separation techniques (liquid
313 chromatography, electrophoresis or mass spectrometry) [80]. Currently, electrochemical glycobiosensors
314 using a lectin as a biorecognition element have been developed to detect changes in the degree of
315 sialylation, fucosylation, galactosylation and mannosylation in cell surfaces and serological samples [81].
316 Overall glycosylation profile of the sample or of a specific glycoprotein can be characterized by
317 electrochemical methods used in glycobiosensors [82, 83].

318 Different electrochemical methods are available for glycobiosensors, including electrochemical
319 impedance spectroscopy (EIS) and voltammetric techniques, such as differential pulse voltammetry (DPV)
320 and square wave voltammetry (SWV) [84, 85, 86]. Among these, EIS is widely used in glycobiosensors as an
321 effective method for detection of lectin-glycan biorecognition on the electrode surface. The measurement is
322 based in the application of a small alternating current amplitude on the electrode interface and subsequent
323 detection of change in impedance during binding events. When the binding event happens, a change in
324 impedance is detected in the presence of a redox probe (eg. electrolyte solution of ferricyanide and
325 ferrocyanide), and the change in electron transfer resistance (RCT) is measured. The change in RCT is due
326 to blocking of charge transfer on the electrode surface in the presence of coating, immobilized biomolecules
327 and biorecognition complex [87]. EIS measurements are transformed in an equivalent circuit and a Nyquist
328 plot to obtain informations about RCT and lectin-glycan binding. By the way, EIS provide a rapid, simple,
329 highly sensitive and label-free detection of glycans in low concentrations, eliminating the use of labels
330 commonly required in non-electrochemical techniques [88]. Voltammetric techniques also have been used

331 for label-free detection of lectin-glycan binding. In general, these techniques are based in the application of a
332 voltage range in the electrode interface, that in the presence of a redox probe, promote a redox reaction and
333 subsequent generation of a current flow [89]. The modification electrode surface with biomolecules and
334 biorecognition complex changes interfacial electron transfer reaction of the redox probe, resulting in changes
335 of current. The variation of current is measured and quantitative information about the glycan target can be
336 obtained. The voltammetric techniques more used for detection of biorecognition events are DPV and SWV
337 due to high sensitivity, quickness (3 to 10 s is necessary for measurement) and minimization of capacitive
338 interference [90].

339 The first electrochemical biosensor using electrode surface modified with lectins was reported by La
340 Belle and coworkers for detection of cancer-associated T antigen (β -D-Gal-[1 \rightarrow 3]-D-GalNAc) [91]. The N-
341 acetyl-galactosamine/galactose specific plant lectin, PNA (peanut agglutinin) was immobilized onto a gold
342 electrode surface previously modified with a mixed self-assembly monolayer, followed by the incubation with
343 nanocrystal CdS-tagged 4-aminophenyl- α -D-galactopyranoside sugar and target sugars (GalNAc, Gal and T
344 antigen) to establish a competitive assay. The lectin-sugar recognition was monitored by square-wave
345 voltammetric stripping current peaks, which decrease with the bind of target glycan. Current peaks decrease
346 for target sugars, in the affinity order T antigen > Gal > GalNAc, achieving a detection of 0.1 μ M for T
347 antigen, 1.0 μ M for Gal and 2.7 μ M for GalNAc.

348 An ultrasensitive label-free glycobiosensor was reported for measurement of sialylated glycoproteins
349 using the EIS method [93]. A sialic acid specific lectin from *Sambucus nigra* (SNA I) was covalently attached
350 to a mixed self-assembled monolayer (SAM) consisting of 11-mercaptopundecanoic acid and 6-
351 mercaptohexanol onto a gold electrode. The biosensor detected glycoproteins fetuin and asialofetuin through
352 the fraction of sialic acid, with a detection limit of 0.33 fM for fetuin and 0.54 fM for asialofetuin. Its potential
353 to detect sialylated glycoproteins with very low limit of detection are attractive for clinical applications in the
354 search of sialylated glycobiomarkers of diseases.

355 Other label-free biosensor based in EIS was developed for detection and discrimination of the cancer
356 glycobiomarker alpha-fetoprotein in serum samples using lectins as biorecognition elements [94]. First, a
357 lectin biosensor for detection of AFP was mounted through the attachment of wheat-germ agglutinin (WGA)
358 lectin to single-wall carbon nanotubes (SWCN) onto a screen-printed carbon electrode (SPCE). The
359 biosensor was used for binding AFP and achieved a detection limit of 0.1 ng/L. Moreover, lectin biosensors
360 were fabricated with different lectins to characterize glycosylation profile of N-glycan linked to AFP and
361 discriminate between healthy and cancer patient's serum samples. The lectins WGA (GlcNAc-specific), LCA
362 (mannose-specific), Con A (mannose-specific), SNA (sialic acid-specific) and DAS (LacNAc-specific) were
363 immobilized onto different SWCN-modified SPCE and changes in charge transfer resistance were monitored.
364 The results suggest an increase of core fucosylation and α 2-6 sialylation in AFP from cancer when
365 compared with healthy serum, being a potential tool for label-free profiling of glycan expression in serum
366 samples to provide early diagnosis.

367 A Con A-based biosensor was reported as a strategy for electrochemical detection of abnormal
368 glycoproteins from serum of patient's infected with dengue virus serotypes 1, 2 and 3 [95]. This biosensor is
369 based on gold electrode modified with phospholipid membrane and Con A to form a lipid-Con A surface,
370 which was incubated with serum from patients. Electrochemical characterization performed by voltammetry
371 and EIS recorded decreases in the current response of the electrodes and increases of charge transfer
372 resistance after incubation with infected sera by dengue virus serotypes 1, 2 and 3. These findings indicate

373 the binding of serum glycoproteins to Con A by its mannose-binding sites, revealing that the biosensor was
374 able to recognize the dengue serotypes and are useful for detection of dengue infections.

375

376 **8 ELECTROCHEMICAL GLYCOBIOSENSORS BASED IN CRAMOLL 1,4 LECTIN**

377

378 Cramoll 1,4 is a legume lectin isolated from *Cratylia mollis* seeds that has been employed as
379 biorecognition element in glycobiosensors. *C. mollis* is a native forage occurring in the semiarid region of
380 Brazil, popularly known as camaratu bean; it belongs to the same Fabaceae family of *Canavalia ensiformis*,
381 the natural source of Con A lectin [7, 88]. The *C. mollis* seeds are sources of four molecular isoforms with
382 diverse specificity denominated Cramoll -1, -2, -3 and -4, and a preparation containing the 1 and 4 isoforms,
383 known Cramoll 1,4. Cramoll 1, 2 and 4 are glucose/mannose lectins, and Cramoll 3 is a galactose specific
384 lectin [7, 89]. Cramoll 1 in a characterization study using X-ray crystallography, showed classic tertiary
385 structure of legume lectins. Its primary structure consists of 236 amino acid residues with 82% homology with
386 Con A structure, and both have identical carbohydrate binding sites to the methyl- α -D-mannopyranose, with
387 which they interact by hydrogen bond even identical, and binding sites for Ca^{2+} and Mn^{2+} metals. The
388 secondary structure is composed exclusively of three β sheets; all connected by turns [67].

389 Cramoll -1 and -1,4 has showed its potential for various biotechnological applications such as
390 antitumor action [90]; mitogenic activity [91]; pro-inflammatory and healing of experimental tissue lesions [92]
391 and anthelmintic [93]. The potential of this lectin to recognize glycans and glycoproteins is a valuable tool
392 for detection changes in glycosylation of diseases through tissues and serum samples. Cramoll 1 showed a
393 higher potential to mark neoplastic mammary tissues [94] and Cramoll 1,4 revealed more intense staining in
394 hyperplasia prostate tissues when compared to prostate carcinoma tissues [1], being a candidate probe for
395 histochemical studies. Other study reported an affinity column of Cramoll 1 coupled to Sepharose CL4B as
396 an efficient matrix to isolate serum glycoproteins, such as lecithin-cholesterol acyltransferase [95].

397 Cramoll 1,4 has been explored in the recognition of serum glycoproteins from patients contaminated
398 with different serotypes of dengue virus using an electrochemical biosensor, a very promising field. Firstly, it
399 was reported the ability of a Cramoll 1,4 biosensor for sensitive detection of glycoproteins in solution [96]. A
400 mix of Cramoll 1,4 with gold nanoparticles (AuNp) and polyvinylbutyral (PVB) was immobilized onto the
401 surface of gold electrodes and the lectin-modified electrode was incubated with solutions containing
402 ovoalbumin. Electrochemical analysis by voltammetric and EIS detected lectin-ovoalbumin interactions in the
403 electrode surface, confirming that immobilization process of Cramoll 1,4 unaffected the recognition sites. In
404 this case, the lectin recognized oligomannose glycans chains linked to ovoalbumin. After, Cramoll 1,4
405 biosensors also were developed to detect dengue serotypes. A gold electrode was modified with Cramoll
406 1,4, Fe_3O_4 nanoparticles and PVB, and applied to evaluate the binding with fetuin in solutions and serum
407 glycoproteins from patients infected with dengue serotypes 1, 2 and 3 (DS1, DS2 and DS3) [8]. EIS and
408 voltametric measurements registered the binding of Cramoll 1,4 with fetuin and serum glycoproteins of DS1,
409 DS2 and DS3, showing a differential response for each serotype and higher interaction with glycoproteins
410 from serum contaminated by DS2. Similar approach was performed to discriminate serum glycoproteins from
411 DS1, DS2 and DS3; also to detect different stages of infection using sera of patients with dengue fever and
412 dengue hemorrhagic fever. An electrochemical biosensor was constructed using a gold electrode, which
413 surface was modified with a nanocomposite of AuNps - polyaniline (PANI) and Cramoll 1,4, and

414 subsequently exposed to different sera [9]. Voltammetric and EIS data showed that Cramoll 1,4 biosensor
415 discriminates the different serotypes and stages of dengue infection. Thus, Cramoll 1,4 is a promisor tool in
416 researches for biosensing of serum glycoprotein profiles associated to diseases and the lectin preparation is
417 a potential candidate for recognition of glycobiomarkers.

418

419 9. CONCLUSION

420

421 Glycosylation is a rich code of informations about physiological and pathological cellular events.
422 Many changes in glycans and glycoprotein profiles are significantly correlated with diseases and their
423 detection provide early diagnostic, staging and prognostic with high sensitivity. When these glycobiomarkers
424 are found in serum samples, they can be measured through techniques more friendly that provide rapid
425 results. Lectins are sugar-binding proteins that recognize alterations in glycosylation profiles and therefore
426 are useful in glycoanalytical techniques, among this, the biosensing. Glycobiosensors based in
427 electrochemical detection is a good strategy for development of simple, rapid, portable and sensitive devices
428 that can detect glycans and glycoproteins in minimal volume of serum samples to search glycobiomakers in
429 clinical diagnostic.

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