


1 Bioelectroanalysis in a Drop: Construction of a Glucose Biosensor

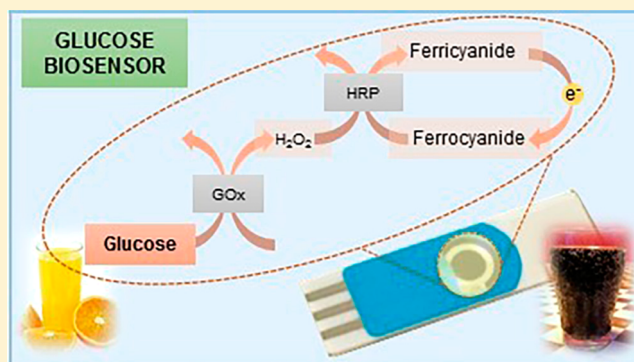
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4  Supporting Information

5 **ABSTRACT:** This lab experiment describes a complete
6 method to fabricate an enzymatic glucose electroanalytical
7 biosensor by students. Using miniaturized and disposable
8 screen-printed electrodes (SPEs), students learn how to use
9 them as transducers and understand the importance SPEs have
10 acquired in sensor development during the last years. Students
11 can also revise concepts related to enzymatic assays, with
12 glucose oxidase and horseradish peroxidase involved in
13 subsequent reactions. Moreover, they learn the trends that
14 current analytical chemistry follows presently such as
15 miniaturization, portability, and low cost. At the same time,
16 this experiment serves to teach basic analytical concepts
17 (accuracy, precision, sensitivity, and selectivity) in a practical
18 way. The high clinical interest of glucose, due to a large number of diabetes patients around the world, and the application of the
19 sensor to analysis of real food samples make this experiment very attractive to students. The questions set out along this
20 experiment help students to acquire skills for solving analytical problems from the very beginning.

21 **KEYWORDS:** Upper-Division Undergraduate, Analytical Chemistry, Laboratory Instruction, Hands-On Learning/Manipulatives,
22 Quantitative Analysis, Electrochemistry, Bioanalytical Chemistry, Biotechnology, Carbohydrates, Enzymes



23 **M**iniaturization is presently one of the most important
24 trends in analytical chemistry. The reduction of the size
25 of analytical systems¹ involves their simplification as well as a
26 decrease in costs, reagents, and sample volume. Furthermore, it
27 is related to many principles of Green Analytical Chemistry.²
28 Electrochemical detection closely connects with these aims
29 because of its inherent ease of miniaturization. Moreover, an
30 improvement in productivity-related properties such as analysis
31 time or cost as well as in others related to environmental
32 benefits like waste production or energy consumption is very
33 advantageous. Other basic analytical properties (e.g., accuracy,
34 precision, sensitivity, and selectivity) are generally not
35 compromised since electrochemical analysis is among the most
36 sensitive detection techniques (as demonstrated by its leading
37 use in biosensing)³ and mass production increases the precision
38 of disposable devices.

39 During the past few years, screen-printing technology has
40 been increasingly used in the fabrication of low-cost thick-film
41 electrodes with small size and good analytical characteristics.
42 During the last years, they have been the basis of many
43 biosensors⁴⁻⁶ because of their low cost and simplicity. Another
44 advantage of screen-printed electrodes (SPEs) is the possibility
45 of doing in situ analysis.⁷ Apart from the electrodes,
46 electrochemical equipment (potentiostats) is also being
47 miniaturized.

48 The aim of this experiment is to build an enzymatic
49 electrochemical biosensor to measure glucose in real food
50 samples. Biosensing is a field that is growing continuously, as
51 demonstrated by the leading place of the journal *Biosensors &*

Bioelectronics.⁸ Glucose is probably one of the most important
52 biological compounds because of its engagement in a multitude
53 of reactions.⁹ Glucose analysis in blood is very important and
54 common because of diabetes mellitus, a disease that is suffered
55 by approximately 150 million people around the world.^{10,11}
56 This disease is produced when the pancreas does not generate
57 enough insulin or when the body cannot use the insulin it
58 produces in an effective way. This leads to an increased level of
59 glucose in the blood. Thus, determination of glucose is one of
60 the most important analytical problems in food science and
61 clinical analysis, so much so that glucose biosensors account for
62 approximately 85% of the entire biosensor market.^{12,13}
63

In this experiment, the students develop an amperometric
64 glucose sensor using the bienzymatic system glucose oxidase
65 (GOx)/horseradish peroxidase (HRP) and ferrocyanide as an
66 electron-transfer mediator.^{14,15} Moreover, since the sensor is
67 fabricated using screen-printed carbon electrodes (SPCEs),
68 students are introduced to the miniaturization of analytical
69 devices.
70

The fabrication of this glucose sensor is based on a very
71 simple procedure reported by our research group,^{14,16,17} and it
72 is addressed to undergraduate students of advanced analytical
73 chemistry. The high educational content related to biosensor
74 principles and new contemporary trends in analytical chemistry
75

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76 also makes this experiment very attractive for the training of
77 chemistry, biotechnology, or biochemistry students.

78 There are many laboratory experiments about detection of
79 glucose using the enzyme GOx, but to the best of our
80 knowledge this is the first undergraduate lab experiment that
81 uses screen-printed electrodes to develop a glucose sensor.
82 Moreover, the combination of a simple procedure with the use
83 of SPCEs eliminates time-consuming maintenance of conven-
84 tional electrodes commonly required with other reported
85 sensors.^{18–20}

86 This laboratory experiment is very useful to introduce
87 students to both electrochemical and biosensor methodologies
88 and provides students with several objectives:

- 89 • Learn the principles of important electrochemical
90 techniques such as cyclic voltammetry and chronoamper-
91 ometry.
- 92 • Use low-cost, disposable, and miniaturized electrodes.
- 93 • Fabricate a glucose biosensor, optimize the parameters
94 influencing the analytical signal, and study the analytical
95 characteristics of the methodology.
- 96 • Analyze real food samples and learn how to validate the
97 methodology.

98 ■ EXPERIMENTAL SECTION

99 This lab experiment is designed for a maximum of 15
100 undergraduate or Master's students working in groups of
101 three during four sessions of 4 h (the lab experiment planning
102 is more detailed in the [Supporting Information](#)). The
103 laboratory experiment consists of the following steps:

- 104 (i) Evaluation of the ferro/ferricyanide system using cyclic
105 voltammetry to set the detection potential.
- 106 (ii) Optimization of the concentrations of enzymes and
107 mediator.
- 108 (iii) Calibration of the biosensor and evaluation of the
109 sensitivity.
- 110 (iv) Study of the precision.
- 111 (v) Evaluation of the selectivity.
- 112 (vi) Determination of glucose in real food samples.

113 Instrumentation

114 Electrochemical measurements were carried out with a
115 μ AUTOLAB potentiostat (Metrohm, Switzerland) interfaced
116 with a computer system and controlled by Autolab GPES 4.9
117 software. Commercial screen-printed carbon electrodes (ref.
118 DRP-110; [Figure 1](#)) and the connector to the potentiostat (ref.

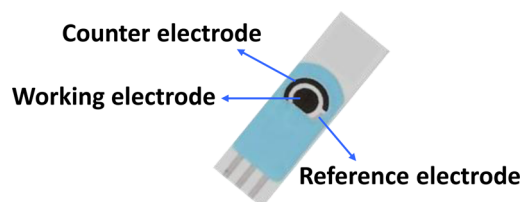


Figure 1. Picture of a screen-printed carbon electrode.

119 DRP-DSC) were purchased from DropSens (Spain). More
120 information on SPEs can be found in the student handout in
121 the [Supporting Information](#).

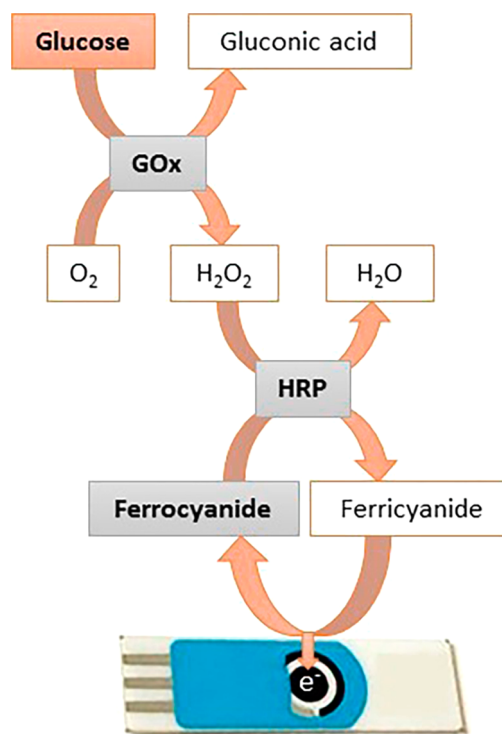
122 Sensor Phase

123 The biosensor constructed in this experiment has a bienzymatic
124 sensor phase consisting of glucose oxidase and horseradish

peroxidase, with ferrocyanide as an electron-transfer mediator. 125
The combination of these enzymes produces an enzymatic 126
cascade of reactions in which GOx and HRP are catalytically 127
linked.²¹ These types of cascade schemes may produce signal 128
amplification and therefore enhance the sensitivity of the 129
biosensor. Another advantage is that by removal of the 130
hydrogen peroxide (H_2O_2) generated, peroxide-induced 131
degradation of the GOx enzyme could be reduced.²² On the 132
other hand, redox mediators are frequently employed in 133
bienzymatic sensors because of their lower detection 134
potentials.^{10,23,24} This is very interesting since it improves the 135
selectivity: at lower potentials (in absolute value), fewer 136
compounds present in the sample are exposed to being 137
oxidized or reduced. 138

In the proposed enzymatic cycle, GOx catalyzes the oxidation 139
of glucose by oxygen, generating gluconic acid and H_2O_2 . Then 140
HRP catalyzes the oxidation of ferrocyanide to ferricyanide, 141
consuming the H_2O_2 previously generated. The analytical signal 142
is the current intensity due to the electrochemical reduction of 143
the enzymatically generated ferricyanide. [Scheme 1](#) shows the 144 s1

Scheme 1. Diagram of the Catalytic Reactions and the Reduction of Ferricyanide Produced on the Electrode Surface, Where GOx, HRP, and Ferrocyanide Are Immobilized^a



^aAdapted with permission from ref 26. Copyright 2016 Elsevier.

reactions involved. Since glucose produces hydrogen peroxide 145
stoichiometrically, and this in turn produces ferricyanide, the 146
concentration of glucose in the measuring solution can be 147
calculated by measuring the concentration of reduced 148
ferricyanide. 149

150 Procedure

Students prepare the biosensors by depositing onto the surface 151
of the working electrode 10 μ L of a mixture containing the 152
enzymes and the mediator at the adequate concentrations, 153

154 prepared in a 0.1 M Tris-HNO₃ buffer (pH 7.0). Then, after a
155 drying step at room temperature (approximately 40–60 min),
156 the sensors are ready to use. If they are going to be employed in
157 the following days or weeks, they must be kept protected from
158 light and at 4 °C.

159 All of the measurements are carried out at room temperature
160 with all three electrodes of the SPCE (working, counter, and
161 pseudoreference) covered with 40 μL of the measuring
162 solution.

163 The analytical signal is the current intensity measured after
164 recording a chronoamperogram (current vs time) at a potential
165 of −0.1 V vs Ag pseudoreference electrode for 100 s (Figure 2).

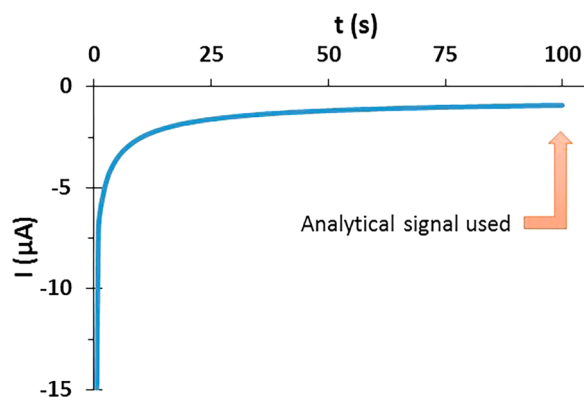


Figure 2. Chronoamperogram obtained at −0.1 V vs Ag pseudoreference electrode, recorded in a 0.5 mM glucose solution with 1.6 units/μL GOx, 2.5 units/μL HRP, and 0.05 M ferrocyanide (in 0.1 M Tris-HNO₃ buffer (pH 7.0)) immobilized on the working electrode (10 μL).

166 A negative current due to the reduction of ferricyanide is
167 obtained, as shown in Figure 2. SPCEs are considered as
168 disposable, and a different sensor is used for each measurement.

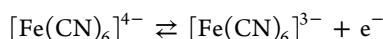
169 ■ HAZARDS

170 Nitric acid, used for the preparation of the Tris-HNO₃ buffer, is
171 corrosive and causes serious burns, so it must be handled with
172 appropriate gloves, safety glasses, and protective clothing. The
173 main hazard of potassium ferrocyanide is that it releases a very
174 toxic gas when it is in contact with acids.

175 ■ RESULTS

176 Electrochemical Behavior of Ferrocyanide

177 After learning about the cascade of enzymatic reactions,
178 students knew that they had to measure the current intensity
179 due to the reduction of ferricyanide. Thus, a potential for
180 ferricyanide reduction had to be applied. Then students
181 investigated the process of the ferro/ferricyanide system,
182 recording a cyclic voltammogram (CV) (one CV can be
183 recorded by each group) in a drop of 0.01 M ferrocyanide
184 solution from −0.2 to 0.8 V at a scan rate of 50 mV/s (Figure
185 3). Previously, a CV was recorded in the background electrolyte
186 to confirm that there was no interference in the potential
187 window scanned. Ferrocyanide shows an electrochemical
188 process according to the following reaction:



189 Looking at the voltammogram, students discussed the
190 reversibility of the process. In this case, the system was

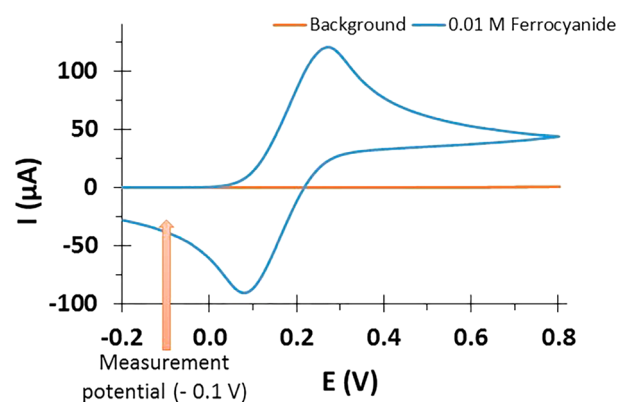


Figure 3. Cyclic voltammogram for 0.1 M Tris-HNO₃ buffer solution (pH 7.0) (background, orange) and for 0.01 M ferrocyanide (blue).

considered to be quasireversible since the difference in
potentials is 183 mV and the ratio of peak currents is 1.1.²⁵

To set the adequate potential for recording the chronoamperograms used for glucose determination, students looked at the process recorded in the CV for the ferrocyanide solution (Figure 3) and chose the potential they considered better (−0.1 V in this case). They argued that at this potential ferricyanide can be reduced to obtain the initial product, ferrocyanide, which is maintained as a complex of Fe(II) on the electrode, and therefore, all of the current intensity comes from the reduction of the ferricyanide (Fe(III)) enzymatically produced. Thus, higher glucose concentrations produce higher current intensities (in absolute value).

Optimization of the Enzyme Concentrations

Students had to know that introducing an analytical methodology requires that once the analytical signal has been identified, the different variables involved must be optimized before the analytical properties (sensitivity, precision, etc.) can be studied. Then students were requested to identify which were the different variables that can influence the signal. After discussion, variables such as pH, electrolyte, and enzyme and mediator concentrations were mentioned. The instructor explained that a 0.1 M Tris-HNO₃ buffer (pH 7.0) was chosen since it was used for similar reported glucose sensors.^{14,26} Thus, the enzyme and ferrocyanide concentrations were identified as relevant variables that should be optimized. With this aim, students prepared sensors with various concentrations of the enzymes, using 10 μL of mixtures with different concentrations of the enzymes and a constant concentration of ferrocyanide. For each mixture, chronoamperograms were recorded in the background electrolyte (buffer solution) and in a 0.4 mM glucose solution.

Figure 4 shows results the students obtained for the different concentrations of enzymes studied. It can be noted that the intensities for glucose solutions were very similar for all of the mixtures, whereas the intensities for the background increased with the concentration of HRP, with the lowest obtained for 2.5 units/μL HRP. For this concentration of HRP, the lowest background was obtained for 1.6 units/μL GOx. Therefore, students chose those enzyme concentrations for the construction of the biosensor.

Optimization of the Ferrocyanide Concentration

The following step was the optimization of the concentration of ferrocyanide. Different concentrations of the electron-transfer mediator (0.05, 0.1, and 0.2 M) were studied using the enzyme

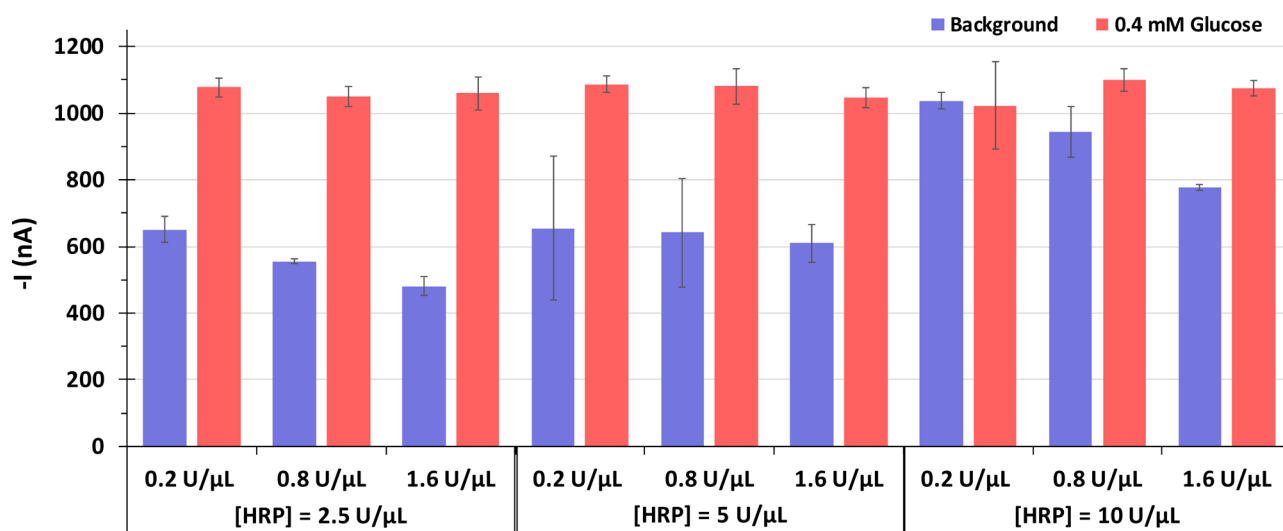


Figure 4. Current intensities recorded at -0.1 V vs Ag pseudoreference electrode after 100 s in buffer solution (background, blue) and a 0.4 mM glucose solution (red). [HRP] = 2.5, 5, or 10 units/ μ L; [GOx] = 0.2, 0.8, or 1.6 units/ μ L; [ferrocyanide] = 0.1 M. Data are given as mean \pm standard deviation (SD) ($n = 3$).

236 concentrations optimized in the previous section (1.6 units/ μ L
237 GOx and 2.5 units/ μ L HRP).

238 **Figure 5** presents the current intensities obtained by the
239 students using different concentrations of ferrocyanide. As can

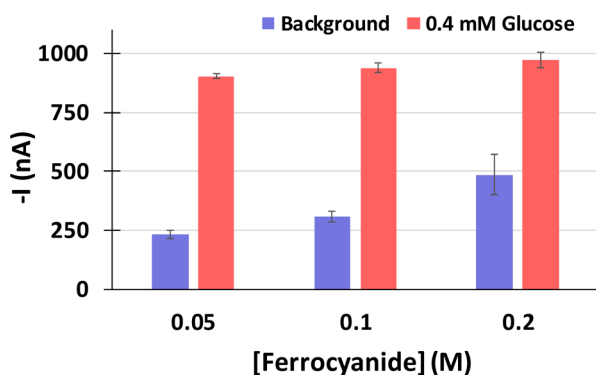


Figure 5. Current intensities recorded at -0.1 V vs Ag pseudoreference electrode after 100 s in buffer solution (background, blue) and 0.4 mM glucose solution (red) with ferrocyanide concentrations of 0.05, 0.1, and 0.2 M using 1.6 units/ μ L GOx and 2.5 units/ μ L HRP (deposition of a 10 μ L drop). Data are given as mean \pm SD ($n = 3$).

240 be seen, the analytical signal for a 0.4 mM glucose solution
241 increased very slightly with the ferrocyanide concentration.
242 However, the intensities in the background decreased when the
243 concentration of the mediator was reduced. Therefore, students
244 concluded that 0.05 M ferrocyanide was the best concentration
245 because it gave a higher signal-to-noise ratio.

246 Calibration of the Biosensor

247 Once the sensor phase had been optimized, students carried
248 out a calibration plot in order to know how the biosensor
249 responded to increasing glucose concentration and to revise
250 some analytical characteristics of the methodology, namely,
251 capital (e.g., accuracy, representativeness), basic (e.g., sensi-
252 tivity, precision), and productivity-related properties (e.g.,
253 analysis time and cost). As shown in **Figure 6**, they found a
254 linear relationship between the current intensity and glucose

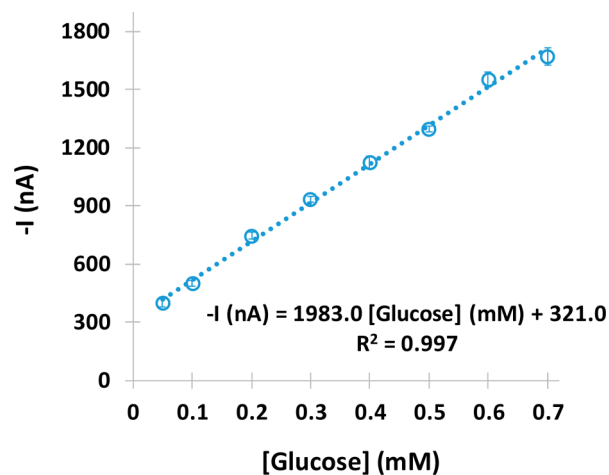


Figure 6. Calibration plot obtained with the glucose biosensor fabricated by immobilizing 10 μ L of 1.6 units/ μ L GOx, 2.5 units/ μ L HRP, and 0.05 M ferrocyanide solution. Data are given as mean \pm SD ($n = 3$).

concentration in the range between 0.05 and 0.7 mM with a
255 sensitivity of 1983.0 nA/mM. Students also calculated
256 important parameters such as the limit of detection (LOD)
257 and the limit of quantification (LOQ) according to the
258 following equations: $LOD = 3s_b/m$ and $LOQ = 10s_b/m$, where
259 m is the slope of the calibration curve and s_b is the standard
260 deviation of the intercept.^{27,28} For this biosensor, the LOD and
261 LOQ values thus calculated were 0.03 and 0.1 mM,
262 respectively. They were also motivated to discuss other glucose
263 sensors found in the literature^{14,22,26,29,30} and to compare their
264 analytical characteristics (sensitivity, precision, linear range,
265 LOD, and LOQ) as well as the procedure of construction
266 (simplicity and fabrication time).
267

268 Precision

The precision of the calibration curve is very important,
269 especially when dealing with an analyte as important as glucose.
270 In order to evaluate it, three calibration curves measured on
271 different days by different groups of students using different
272 solutions were compared (data are shown in **Table 1**). They
273 11

Table 1. Equations of the Calibration Plots for Three Glucose Biosensor Series

Calibration Curve	Slope, nA/mM ^a	Intercept, nA ^b	R ²	Linear Range, mM
1	1983.0	321.0	0.996	0.05–0.70
2	2136.0	168.0	0.997	0.05–0.70
3	2105.0	230.0	0.991	0.05–0.70

^aMean \pm SD = 2080 \pm 81. ^bMean \pm SD = 240 \pm 77.

274 showed very good reproducibility, with a relative standard
275 deviation of the slopes of 3.9% ($n = 3$). In this way, students
276 understood that a biosensor must present good reproducibility
277 notwithstanding the fabrication day, the day of use, and the
278 operator. Since the temperature is not controlled (experiments
279 were done at room temperature), this was also indicative of the
280 robustness of the methodology.

281 Selectivity

282 Selectivity is another important property of a biosensor that has
283 to be taken into account. Students evaluated how the presence
284 of some species affected the analytical signal. In this case,
285 fructose and ascorbic acid were chosen as possible interferences
286 that could be found in real samples. Thus, different biosensors
287 were constructed for determination in mixtures of glucose/
288 fructose and glucose/ascorbic acid. The results obtained are
289 reported in Table 2.

Table 2. Study of Fructose and Ascorbic Acid Interferences in the Glucose Sensor

Sample	-I, nA	\pm SD, nA ^b
Background	386	11
Glucose 0.3 mM	791	21
Glucose 0.3 mM/Fructose 0.3 mM	852	30
Glucose 0.3 mM/Ascorbic Acid 0.3 mM	538	25
Glucose 0.3 mM/Ascorbic Acid 6 μ M ^a	829	29

^aRatio in real samples. ^b $n = 3$.

290 As can be seen in Table 2, fructose and ascorbic acid
291 produced the opposite effect on the analytical signal. In the case
292 of fructose, a slight increase in the signal was seen; meanwhile,
293 ascorbic acid (usually employed as an antioxidant) produced a
294 decrease in the signal. The effect is not so important when the
295 glucose:ascorbic acid ratio is similar to that found in real
296 samples (e.g., orange juice³¹). Students were encouraged to
297 look for possible solutions to avoid the interference produced
298 by ascorbic acid and incorporated their ideas in the lab report.
299 Students indicated as a good idea coating the sensor with a
300 Nafion film since it is a negatively charged polymer that repels
301 anions.^{32,33}

302 Application to Real Samples

303 The final aim of the biosensor developed here was to determine
304 glucose concentrations in real samples. Thus, students
305 determined glucose concentrations in a cola beverage and
306 orange juice purchased in the market. The only pretreatment
307 needed was dilution of the sample to obtain a concentration
308 within the linear range of the biosensor (dilutions were made in
309 0.1 M Tris-HNO₃ buffer solution, pH 7.0). Previous degassing
310 by stirring was required for cola samples.

311 The samples were validated previously by the instructor
312 using an alternative method (a commercial glucose enzymatic
313 assay kit with spectrophotometric detection), and the results

were given to the students after they analyzed the samples. In
this way, students compared the results obtained using the
biosensor with the results given by a “reference method”. The
values for glucose concentration obtained with the sensor and
the commercial kit are summarized in Table 3. Students

Table 3. Application of the Biosensor to Analysis of Real Samples

Sample	Glucose Concentration Determination, g/100 mL ^a	
	Electrochemical Sensor	Spectrophotometric Assay
Cola beverage	3.3 \pm 0.3	3.42 \pm 0.03
Orange juice	3.4 \pm 0.2	3.47 \pm 0.04

^aData are given as mean \pm SD calculated with two degrees of freedom and $p = 0.05$; $n = 3$.

statistically compared the mean values obtained using the two
methodologies through a Student's t test.³⁴ The t values
calculated for the cola beverage and orange juice were less than
the t value tabulated for two degrees of freedom and a 0.05
significance level. Thus, there were no significant differences
between the glucose concentrations given by the biosensor and
the enzymatic assay.

DISCUSSION

After constructing the glucose biosensor, the students learned
about different electrochemical techniques and their application
to the development of an enzymatic biosensor. They also
realized that the use of low-cost, disposable, miniaturized, and
portable equipment is of paramount importance today,
especially when real samples are analyzed.

The experiment was completed with discussions on the
following topics:

- (i) The analytical problem: types of samples and levels of glucose.
- (ii) The state of the art: previous works obtained from a bibliographic search on glucose electrochemical enzymatic sensors, discussing also the different generations of sensors, the role of nanomaterials, and nonenzymatic approaches.
- (iii) Evolution of electroanalysis from conventional cells to miniaturized designs.
- (iv) Analytical properties (accuracy, precision, sensitivity, selectivity, and especially those related to productivity: analysis of time and cost—see the Supporting Information) and approaches for improving them.

CONCLUSIONS

This experiment served as a practical introduction to biosensor technology and to the challenge of glucose determination. The high number of diabetes patients worldwide increased its relevance, and the application of the developed sensor in analysis of real samples stimulated the students' interest. This lab experiment also introduced students to the miniaturization and simplification of analytical devices and methodologies, some of the most important trends in modern analytical chemistry. Biosensors are an excellent example of simple and promising analytical tools, and students could become familiarized with their two components (sensing zone and transducer), understanding concepts of enzymatic analysis and electroanalysis at the same time. Moreover, they learned how to use screen-printed electrodes, which are widely used today in

363 the development of sensors. They also discussed analytical
364 properties (e.g., accuracy, precision, sensitivity, and selectivity)
365 and productivity-related features (e.g., analysis time and cost).
366 In summary, this lab experiment allowed students to acquire
367 problem-solving skills, to reach a high level of critical thought,
368 and to be more confident in facing real-world analytical
369 problems.

370 ■ ASSOCIATED CONTENT

371 ● Supporting Information

372 The Supporting Information is available on the ACS
373 Publications website at DOI: [10.1021/acs.jchemed.6b00948](https://doi.org/10.1021/acs.jchemed.6b00948).

374 Notes for instructors and a suggested student handout
375 ([PDF](#), [DOCX](#))

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382 Notes

383 The authors declare no competing financial interest.

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