Graphene electrochemical sensors for the detection of S 29:1-(2-chloro-2-(4-chlorophenyl)ethyl)-N-(4-fluorobenzyl)-1H-pyrazolo[3,4-d]pyrimidin-4-amine: an anticancer drug

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Abstract

Graphene (G) and Graphene derivatives (Gds) represent the next generation of smart materials, having exciting features to apply in Medicine field. Especially, several studies [1,2] report about the biocompatibility of graphene and graphene derivatives in several different cell lines. Biocompatibility for G and Gds seems to be related to main aspects, as: the metal-free chemical composition of graphene and the nano sheets shape. Recently, Valentini, et al. [3] report about the in vitro biocompatibility of Graphene Oxide (GO) nanosheets, tested in different human cell lines and compared with that exhibited by Single Wall Carbon Nanotubes (SWCNTs).

This GO nanomaterial was obtained during 2015 in Tor Vergata University by the Patent N102015000023739, applying the electrochemical exfoliation of highly oriented pyrolitic graphite anode electrodes, in electrolytic working medium. The chemical-physical features and also the biocompatibility of GO sheets, synthesized in this work, were widely reported in our previous paper [3]. Furthermore, antibacterial properties were studied, in presence of Staphylococcus Aureus and Escherichia Coli bacteria, as reported in our previous paper [4].

In this paper, the electrochemical behavior of graphene was investigated in presence of 1-(2-chloro-2-(4-chlorophenyl)ethyl)-N-(4-fluorobenzyl)-1H-pyrazolo[3,4-d]pyrimidin-4-amine (labeled as S 29 and, see Figure 1), for the first time. The scientific interest toward this therapeutics was mainly due to the inhibitor role toward the sarcoma (Src) related protein kinase, especially in neuroblastoma, medulloblastoma and osteosarcoma cancer cells [5-7].

For this purpose, the great possibility to assemble an electrochemical portable sensor, able to quantitatively detect S29, during the inhibitor processes in neuroblastoma cell lines, seems to be an extraordinary challenge, in the anticancer drug research fields. According to this consideration, the aim of this study is to assemble and then characterize new miniaturized graphene modified Screen Printed Electrodes (SPEs) system, combined with an electrochemical detection for the quantification of S29 therapeutics. This latter was kindly provided by Prof. M. Botta, and here, a short description of the S29 synthesis was also provided. The synthetic approach to prepare the compound S29 [8] was started from the same reaction between ethylethoxymethylene-

**Figure 1.** S 29 structure: 1-(2-chloro-2-(4-chlorophenyl)ethyl)-N-(4-fluorobenzyl)-1H-pyrazolo[3,4-d]pyrimidin-4-amine
in CHCl₃ led to the formation of the halogenated derivative [11]. The C₄ chlorine atom was in turn substituted with (4-fluorophenyl) methanamine, in anhydrous toluene at room temperature, to afford the final compound in yields of 75%.

The electrochemical profile of S29 was investigated by applying GO modified Screen Printed Electrodes (GO/SPEs). SPEs are commercially available (purchased by Metrohm DropSens, Origgio (VA) Italy) and then, they are modified by drop casting 6μL of GO nano dispersion, with a concentration of 1mg/mL, in ethanol. The electrochemical characterization study was performed using these modified GO Screen Printed Electrodes, which correspond to the electrochemical cell, incorporating three electrodes in the same system: a GO based working electrode (3-mm diameter), a Ag/AgCl reference electrode, and a carbon pseudo-counter electrode. An Autolab electrochemical system (Eco Chemie, Utrecht, The Netherlands) equipped with PGSTAT-12 and GPEPS software (Eco Chemie, Utrecht, The Netherlands) was applied for the electrochemical investigation of the electron-transfer properties, exhibited by S29 on GO/SPEs, mainly by using Differential Pulse Voltammetry (DPV) techniques. The electrochemical oxidation profile for S29 was reported on Figure 2A, with an oxidation peak centered around +400 mV, versus Ag/AgCl reference electrode. The corresponding calibration curve was also carried out and reported on Figure 2B and C. The electrochemical performances of the amperometric sensor seem to be relevant, showing a wide range of concentration (10-10000 nmol/L), a low detection of limit (5 nmol/L), a high sensitivity (0.07 μA cm⁻² nM⁻¹), good reproducibility (~1%) and short response time (4s). This sensor resulted very innovative because in literature, there is only one scientific paper [9], where Martic, et al. reported on an electrochemical biosensor developed for the determination of sarcoma (Src)-related protein kinase, in presence of pyrimidine-based small molecules, as: pyrazolopyrimidines 4-amino-5-(4-methylphenyl)-7-(tert-butyl)pyrazolo pyrimidine (PP1), 4-amino-5-(4-chlorophenyl)-7-(tert-butyl)pyrazolo pyridine (PP2), which are the standard inhibitors for kinases belonging to the Src family of protein; and the inhibitor of Src protein kinase, 2-oxo-3-(4,5,6,7-tetrahydro-1H-indol-2-ylmethylene)-2,3-dihydro-1H-indole-5-sulfonic acid dimethylamide (SU6656). The electrochemical principle of this biosensor was completely different than the GO/SPEs, presented in this study and in any case, the analytical performances of the biosensors, prepared by Martic, et al. [9] were not comparable with those observed in this paper (working with GO/SPE systems). Indeed, the linear range of concentration (5-30 nmol/L) resulted much more limited, working with the inhibition based electrochemical biosensors; with even worse reproducibility (RSD% 3-5) of the measurements and a longer response time (20 min).

In addition, the electrochemical GO/SPEs were highly selective sensors, mainly toward one of several isomeric forms of pyrazolopyrimidines, as the pyrazolo[3,4-d] pyrimidine (labeled as SI 306) (Figure 3). This latter was also kindly provided by Prof M. Botta (Siena University) and synthesised according to the literature [10]. The high selectivity, exhibited by the GO/SPE devices, certainly could represent a future perspective in Medicine, providing the great opportunity to specifically quantify both anticancer drugs (as S29 and SI306).

In conclusion, in this work GO was electrochemically synthesised and fully characterized, for the assembly of modified SPEs, applied for the quantification of S29 therapeutic molecules. Analytical results showed a wide range of concentration, a low detection of limit, good sensitivity, high reproducibility and very short response time. This means that the innovative SPE/GO tools are suitable for in real time pharmacokinetic studies, using disposable, miniaturized and partially automatized electrochemical tools.

Further perspectives and interesting challenges in Medicine, will be represented by the Wireless transmission of the analytical signals (also combined with the IoT_Internet of Things, and ICT_information Comunication Technologies) and the remote control of big data systems.

References

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