Tailoring Molecular Imprinted Polymers as Biomimetic Recognition Coatings for Anti-Diabetic Drugs

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Abstract:
Molecular recognition of pharmaceutical molecules in complex mixtures is a subject of significant interest both for clinical analysis and routine quality control assays in pharma industries. The ever increasing use of anti-diabetic drugs e.g. sitagliptin and metformin also demand their selective extraction and quantitative determination. In present study, we tailored molecular imprinted polymer matrix as highly responsive chemical sensor coatings for molecular recognition of above mentioned anti-diabetic drugs. Molecular imprinted coatings were synthesized as poly(methacrylic acid) cross-linked with ethyleneglycol dimethacrylate following free radical initiated polymerization. The imprinted polymer was processed in gel form and integrated with interdigital capacitors (IDCs) as thin layers by spin coating. The prime advantages of using IDCs are their low fabrication cost, rapid response time and ruggedness. The synthesized imprinted sensor layers have shown appreciable response for target analytes comparing to non-imprinted/control material. Therefore, the developed sensor coatings can be suitably used for specific drug analysis and related bioassay of anti-diabetic drugs.

Key words: Molecular imprinting, anti-diabetic drugs, IDCs, chemical sensors, sitagliptin, metformin.

Introduction
The binary formulations of sitagliptin and metformin have shown suitable potential for treating type 2 diabetes. As there is a growing interest in using these drugs for combination therapy since they provides adequate glycemic control [1]. In this perspective, analytical protocols dealing with selective recognition of these drugs in complex mixtures are essentially required for their quantitative estimation.

Molecular imprinting is a well known method for synthesizing synthetic recognition materials [2] having predefined functionality. Aside from typical applications of imprinted polymers, the prime interest in using them is the flexibility in their processing e.g. in micro/nanoparticles as well as thin layers. This allows using molecular imprinted polymers (MIPs) for separation and sensing purposes, respectively. In view of sitagliptin and metformin presence in complex pharmaceutical formulations and biological samples herein, we developed molecularly imprinted materials as highly responsive sensor coatings combined with interdigital capacitors (IDCs) [3] for anti-diabetic drugs.

Experimental
We prepared molecular imprinted polymer using methacrylic acid (MAA) as functional monomer along with ethyleneglycol dimethacrylate (EGDMA) as cross linker in dimethylformamide (DMF) as solvent. This mixture was heated for 30 minutes at 65°C to form pre-polymer complex. After that, azoisobutyronitrile (AIBN) was introduced to initiate free radical polymerization and heating was continued for another 15 minutes. The resultant imprinted polymer was processed as viscous gel which was diluted accordingly for coating IDCs. The comb shaped interdigital electrode pattern was developed by screen printing. The non-imprinted gel was synthesized under the same conditions except adding template (sitagliptin/metformin). The synthesized polymer gels were developed on IDCs as thin layers by spin coating at 2000rpm. All the sensor measurements were conducted by connecting coated IDCs with Sourcetronic ST2817B LCR meter.
Discussion
Molecular imprinted materials can be tailored according to desired application for instance; they can be processed into sorbent particles as well as sensor coatings. A schematic diagram of synthesizing MIP in gel form has been shown as follows.

In previous studies [4], we showed that sitagliptin or metformin imprinted particles could be used efficiently for respective drug rebinding. The cross sensitivity of the prepared imprinted particles was substantially high for target drug rebinding. Now, we extended this strategy and processed imprinted polymers as thin layers coated on IDCs. For sensor measurements, the IDCs were connected to LCR meter. On exposing to standard drug concentration, a shift in capacitance was observed which was taken as sensor response. For instance, figure 1 shows the capacitance shifts for sitagliptin-imprinted and non-imprinted layers exposed to 10 ppm sitagliptin standard solution.

Conclusion
Molecular imprinted particles have already shown their potential for extraction of anti-diabetic drugs. In this work, we demonstrated that imprinted polymer can be processed as thin films and by combining with IDCs they offer appreciably high sensor response for target pharmaceutical molecules. Additionally, the layer-analyte interactions are reversible which allows reusing sensor interface for several measurements. The use of IDCs as sensing transducer is favorable for obtaining rapid shifts in sensor response and for being low cost they can be fabricated for developing viable sensor devices.

References