

PREPARATION AND ANALYSIS OF MOLECULARLY IMPRINTED SOLID-PHASE EXTRACTION FOR DIAZEPAM AND ITS PHARMACEUTICAL APPLICATIONS BASED ON ACID MONOMERS USING A BULK POLYMERIZATION METHOD

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ABSTRACT

A molecularly imprinted polymer was used to make two electrodes (MIP). MIP was manufactured using Diazepam (DZP) as the template, 2-acrylamido-2-methyl-1-propan sulfanic acid as the monomer, and N.N-methylene bis acrylamide as the cross-linker. Non-imprinting polymers (NIP) were created using the identical composition, minus the template (Diazepam). Tritolyl phosphate (TTP) and dibutyl sebacate (DBS) are examples of plasticisers utilised in the PVC matrix to produce films. DZP-MIP electrode slope, detection limit, durability, and linearity range are evaluated. The outcomes of the selectivity measurements on the interfering cations (Al^{+3} , Ca^{+2} , and K^{+1}) indicate that they do not inhibit Diazepam. The produced electrode exhibited favorable response, including to conduct research on pharmaceuticals.

Keywords: Molecularly imprinted electrode, diazepam, potential metering, 2-acrylamido-2-methyl-1-propan sulfanic acid monomer, and different plasticizers (DBS) (TPH).

تحضير وتحليل الطور الصلب المستخلص المطبوع جزيئياً للديازيبام وتطبيقاته الصيدلانية المعتمدة على المونمرات الحامضية باستخدام طريقة البلمرة الثقيلة

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الخلاصة

تم استخدام بوليمر مطبوع جزيئياً لصنع قطبين (MIP). تم تصنيع MIP باستخدام Diazepam (DZP) كقالب، 2-acrylamido-2-methyl-1-propan sulfanic acid as monomer و N.N-methylene bis acrylamide باعتباره الرابط المتقاطع. تم إنشاء البوليمرات غير المطبوعة (NIP) باستخدام تركيبة متطابقة، باستثناء القالب (Diazepam). يعتبر تريثلول فوسفات (TTP) وثنائي بيوتيل سيبيكات (DBS) أمثلة على الملدنات المستخدمة في مصفوفة PVC لإنتاج الأغشية. يتم تقييم منحدر القطب Dia-MIP، وحد الكشف، والمتانة، والمدى الخطي. تشير نتائج قياسات الانتقائية على الكاتيونات المتداخلة (K^{+1} و Ca^{+2} و Al^{+3}) إلى أنها لا تثبط الديازيبام. أظهر القطب الناتج استجابة إيجابية، بما في ذلك إجراء البحوث على المستحضرات الصيدلانية. على المستحضرات الصيدلانية.

الكلمات المفتاحية: قطب كهربائي مطبوع جزيئياً، ديازيبام، قياس الجهد، مونومر 2-acrylamido-2-methyl-1-propan sulfanic acid، والملدنات المختلفة (DBS) (TTP).

INTRODUCTION

Diazepam, 7chloro1, 3dihydro1methyl5phenyl2H1, 4benzodiazepin2one, is the most often prescribed benzodiazepine hypnotic, tranquilliser, anticonvulsant, and muscle relaxant (Hosseini & Motaharian, 2015). Rapid and dependable screening procedures for drugs and poisons in highly intricate biological specimens (urine, serum) for use in forensic and clinical toxicology (Liu *et al.*, 2013). Diazepam is one of the most often prescribed 1,4-benzodiazepines and is commonly marketed under Valium (Honeychurch *et al.*, 2013).

The effects of sleeping aids. DIA is a benzodiazepine that provides a positive allosteric regulation receptor for gamma-aminobutyric acid, increasing receptor-binding GABA molecules. This modification will induce repolarisation of GABA receptors (channel ligand) and have a calming effect (Omar & Eesa, 2017). The receptors are located in the central nervous system and the mechanism explanation by which DZP induces drowsiness and reduces stress. In Indonesia, DZP is considered a substance class of psychoactive medications, and its prescription is subject to government regulation. To obtain DZP, one must visit a physician and get a prescription (Hasanah *et al.*, 2021).

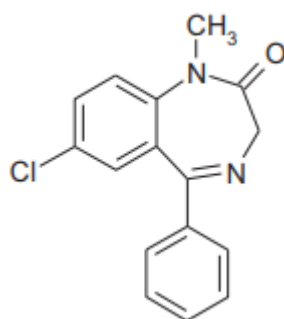


Figure (1): Chemical structure of Diazepam (Casarrubea *et al.*, 2012).

Experimental

Preparation of MIP and NIP

- To prepare Diazepam molecularly imprinted polymer (DZP-MIP1), (0.2847g) of Diazepam was combined with 1 ml of 2-acrylamido-2-methyl-1-propan sulfanic acid as the monomer. Next, 3.08 g of N.N-methylene bis acrylamide was included in the mix as the cross-linker, and (0.3g) of benzoyl peroxide was added as the initiator. The amalgamation was agitated for 5 minutes to achieve a homogenous solution, and then oxygen was removed from the mixture by passing N₂ over it for 20 minutes. The resulting tube was inserted. a container filled with water at 65°C. The molecularly imprinted polymer resulted as soon as the procedure ended completely solidified, and following the polymerisation process, it disintegrated into a tiny polymer particle. Sonication was used on this material. in a 9:1 mixture of CH₃OH and CH₃COOH, eliminating the MIP sample document. How big are DZP-MIP particles (75-125 m) non-molecularly imprinted polymers can be made in the same way components and under the same circumstances as molecularly imprinted polymers. DZP-MIP, but without the Diazepam (template).



Instruments

This study utilised a WTW model ion analyser, a WTW model pH 720 pH metre, and a calomel electrode that has reached saturation (Gallenkamp, USA). The BHH-MIP was an electrode fabricated in a test tube, and every potentiometric experiment was performed at room temperature. In conjunction with the Ag-AgCl and the reference electrode, the diazepam-MIP electrode was a 0.1 M diazepam dissolved in the internal fluid. Placing the PVC tube (1-4 cm in length) in a clear dish and soaking it in THF caused it to become flattened and polished. A membrane was trimmed to match the outer diameter of the PVC tubing and adhered to the precise result. The opposite orientation of the PVC and, finally, the tubing was fastened towards the electrode device. The electrodes were enhanced by good soaking them in 0.1 M diazepam solution for a minimum of three hours prior to use.

Materials and chemicals

Diazepam was acquired to do with the State Corporation for Pills Industry and Healthcare Equipment (IRAQ- SID- Samara). Commercial diazepam pills bought from nearby pharmacies include VALIAPAM 10 tablets 500 mg from (SDI-Iraq), Valium 10 tablets 5 mg from (Australian), and. Plasticizers, Tritoly phosphate 90% was utilised as received and dibutyl sebacate (DBS) (purity of 97.0%), were acquired from Sigma-Aldrich. As a monomer, allyl chloride was used; ethylene glycol di methacrylate (EGDMA) and benzoyl peroxide (BPO)(78%) were acquired from Sigma-Aldrich. The chemicals utilised were of the purest concentration of reagents and were used without extra cleansing.

The creation of standard solutions

Producing a common fluid of 0.01 M Diazepam by breaking down 0.1423 g of legal Diazepam in methanol and then filling a 50 mL volumetric flask with the resulting solution. Using the same method, the additional solutions were produced in 50 mL at concentrations ranging from (5×10^{-5} to 10^{-2}) M. All competing cations (Al^{+3} , Ca^{+2} , and K^{+1}) were made as a 0.01 M stock solution at concentrations ranging from (5×10^{-5} to 10^{-2}) M and then reduction of to 100 ml.

Synthesis of Membrane Molecularly Imprinted Polymers Electrode

According to Thomas and Moody (Moody & Thomas, 1988) , the Diazepam membrane was immobilised within the PVC tube (17) DZP-MIP (0.036g) was combined with other plasticisers (0.45g) employed in this study, including TTP(electrode M1), DBS, and others (electrode M2). Then, 0.2g of It was PVC powder. Added 7 mL of tetra hydro furan and mixed until a thick, viscous liquid was obtained. The fluids were then stirred until the combination was complete and homogenous. The whole combination was poured into a glass ring (30-35 mm in diameter) and set on a flat glass with a superimposed ribbon filter. At ambient temperature, the solvent was permitted to vaporise for at least (24-48) hours. The obtained membrane had a varied thickness than other membranes, ranging between (0.4) and (0.7) mm. This membrane size was appropriate for preparing electrodes.

Construction of Ion-Selective Electrodes

Electrode body structure and immobilisation were accomplished as Mahajan *et al* (Mahajan & Sood, 2007). described. The glass tube had been filled. With 0.1 M diazepam fluid as an internal fluid. Membrane electrode requirements include soaking the membrane in a standard (0.1) M diazepam solution for at the very least two to three hours prior actual metrics (Aljabari & Al-Bayati, 2023).

Pharmaceutical Sample Preparation

Extract the active ingredients from pharmaceutical samples by grinding the crushing tablets using a mortar and pestle. Then, take an acceptable quantity required in advance for use in 50 mL of solutions. Utilised a correct methanol concentration (CH₃OH) for dissolving pharmaceutical samples and filled the volumetric flask to 50 mL with methanol while stirring for more than 30 minutes. After filtering the fluid via 0.07m cellulose filter paper, amounts or proportions of 5×10^{-3} M and 5×10^{-4} M diazepam were achieved.

3. Results and Discussion

The polymer molecular imprint of Diazepam was identified after it was determined using UV. Via the drug's wavelength. As demonstrated in the figures, a preliminary diagnosis was made to confirm the presence of the drug in this imprint.

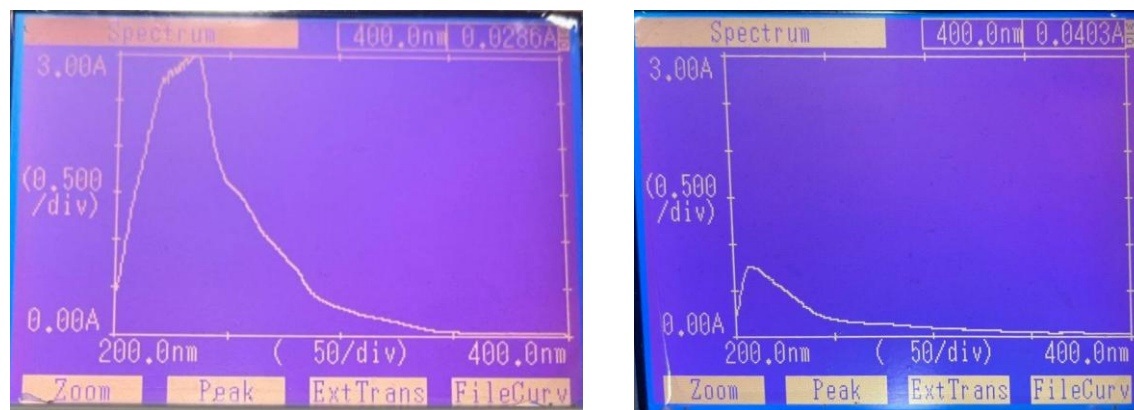


Figure (2): A and B the absorption of diazepam drug at 235.5 nm before and after extraction . MIP of Ate was synthesised using bulk polymerisation (non-covalent). A functional monomer played a crucial role in researching interactions with the template. The MIP and NIP were made using allyl chloride as the monomer. FTIR analysis FTIR is an essential chemical characterisation technique for detecting functional groups in a molecule.

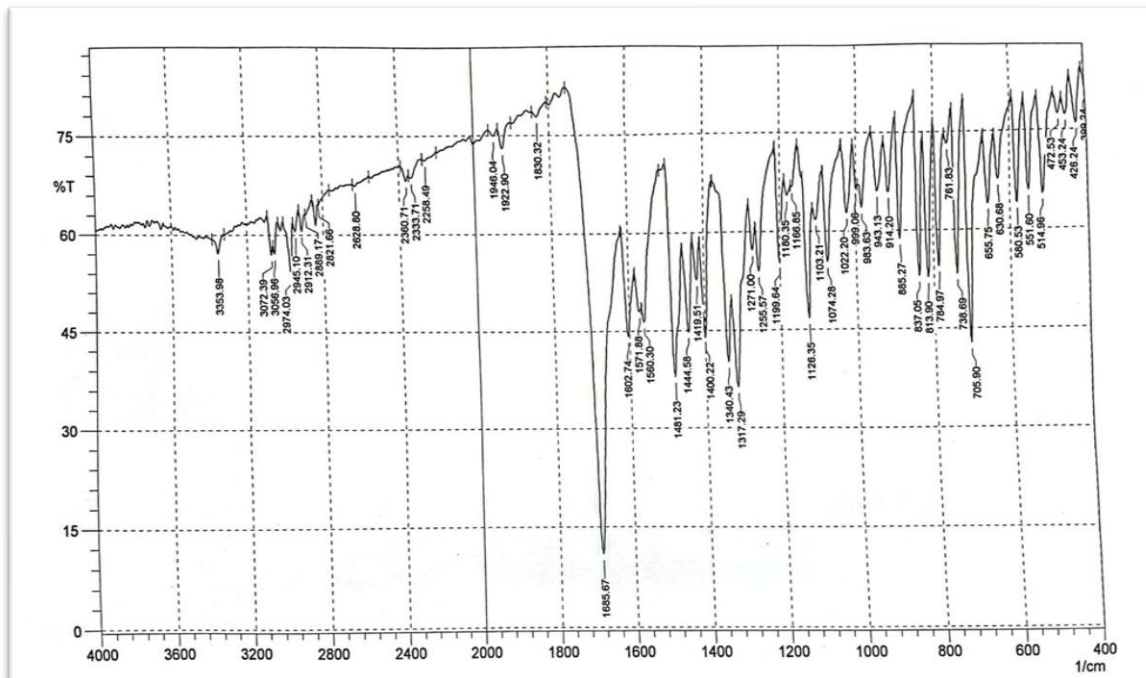


Figure (3): FTIR spectrum of diazepam standard.

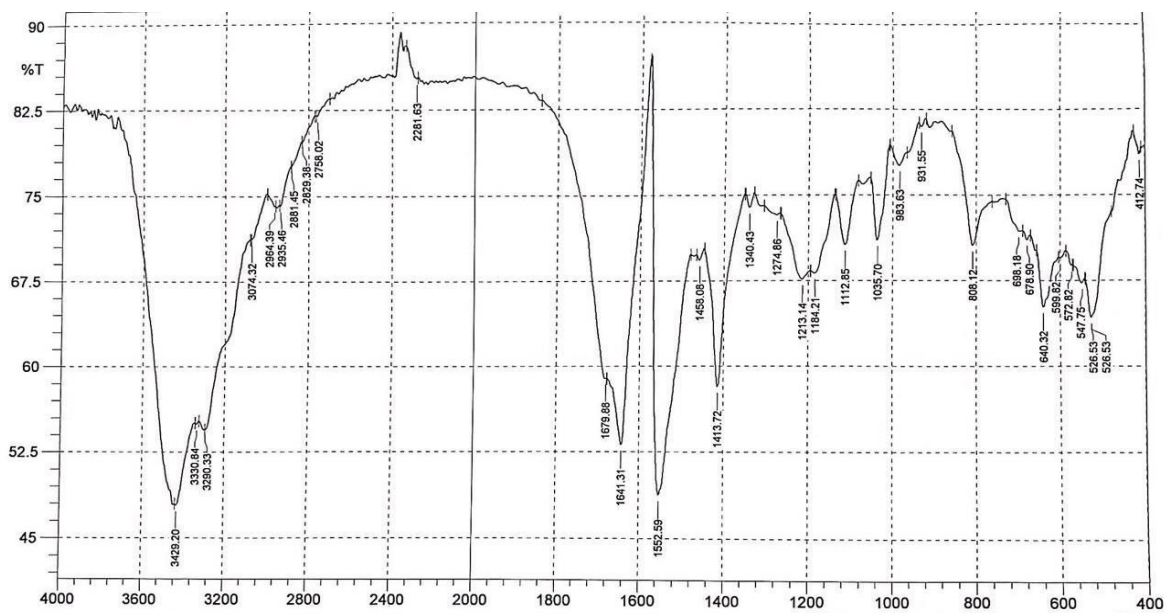


Figure (4): FTIR spectrum of DZP-MIP prior to and following elimination (before removal of the template Diazepam).

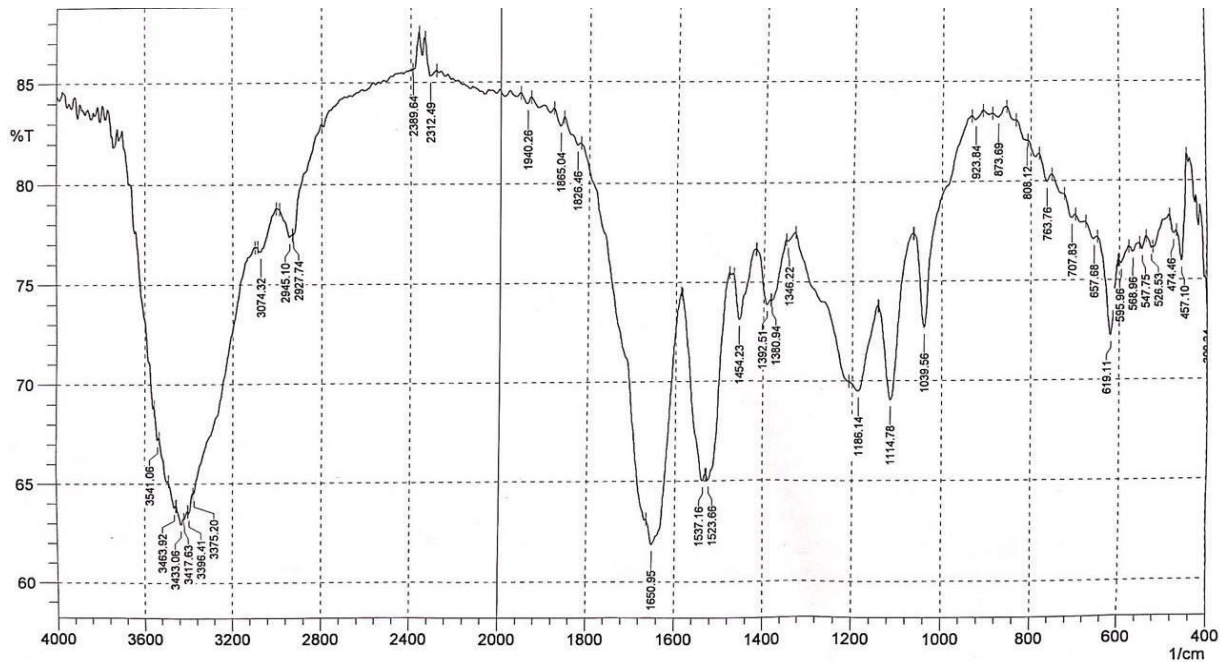


Figure (5): FTIR spectrum of DZP-MIP prior to and following elimination (after removal of the template Diazepam).

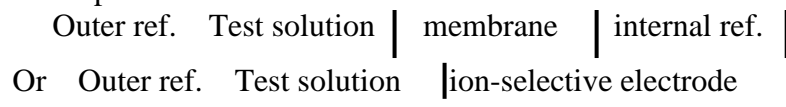
Table (1): The structures of the main three compositions of Dia-MIP and the bands indicate MIP before & after the removal template.

Template (Diaspam)	Monomer (Allyl chloride)	Cross linker (Ethylene glycol di amethacrylat)	
Band	Drug(Template)	MIP before extraction	MIP after extraction
N C-H aliph. str.	2945,2912cm ⁻¹	2964,2935 cm ⁻¹	-
N C-H aromatic str.	3056 cm ⁻¹	3074 cm ⁻¹	2945,2927 cm ⁻¹
N O=C-N str. amid	1685 cm ⁻¹	1641 cm ⁻¹	-
N C=C aromatic. str.	1602 cm ⁻¹	1552 cm ⁻¹	-
N C-Cl	813 cm ⁻¹	808 cm ⁻¹	-
NN-H str.	-	3330 cm ⁻¹	3396 cm ⁻¹
N-S=O str.	-	1340 cm ⁻¹	1380 cm ⁻¹
O-S-OH str.	-	3424,3074 cm ⁻¹	3417,3433 cm ⁻¹
N-C=O carbonyl.	-	1679 cm ⁻¹	1650 cm ⁻¹

The FTIR spectra of Ate displayed the following bands: (2945, 2912, 3056, 1685, 1602, 813) cm⁻¹ for NC-H aliph. str, NC-H aromatic str.,N O=C-N str. amid, NC=C.aromatic. Str. N-C-CL str. N N-C=O carbonyl. Before template removal, the FTIR spectra of the Diazepam MIP(DZP) displayed the following bands: 2964, 2935, and 1641 cm⁻¹ for N C-H alph str., NO=C-N str. amid, and N C=C aromatic str. The absence of N C-H alph str, NO=C-N amid str, and N C=C aromatic str in the FTIR spectrum of the MIP(DZP), following template removal indicates that the drug has been extracted from the template. Using allyl chloride as a monomer for synthesising other MIPs for Diazepam (DZP), the FTIR spectra of the MIPs before and

after removing the template and NIP are shown in Table. The values of the band (Abbas *et al.*, 2020) Ion-selective electrodes (ISE) are among the most widely used types of band electrodes. commonplace sensors based on voltage analysis. Utilised in laboratory experiments, industrial applications, process control, physiological assessments, and ecological monitoring. Membranes of electrodes that respond to Reaction analysis of concentration that generate ions that may be detected using an ion-selective electrode (Aljabari & Al-Bayati, 2021). Electrodes with a membrane are divided into two primary categories: ion-selective, which are sensitive to ions, and molecular-selective, which is sensitive to molecules which are utilised for measuring molecular analytes (Al-Nisani *et al.*, 2021, Ameen *et al.*, 2015). Electronic current travels along electrons in metals but along ions in liquids. Ion-selective electrodes operate based on these two distinct forms of electrical conductivity (Ismaeel & Al-Bayati, 2021). It is possible to do electrical analysis using one of these galvanic cells, electrolysis, and used to assess the conductivity of each electrochemical reaction (Muhammad, 2013).

These cells must be in touch, having fluid across the membrane and inside the cell. There are further ISE configurations where wires are attached to the membrane on only one side. Conventional cell composition consists of:



a device's internal electric current An electrolytic cell is required equal zero. Based on this need, the cell is created based on the terms of the underlying principle behind the construction of electrolytic cells.

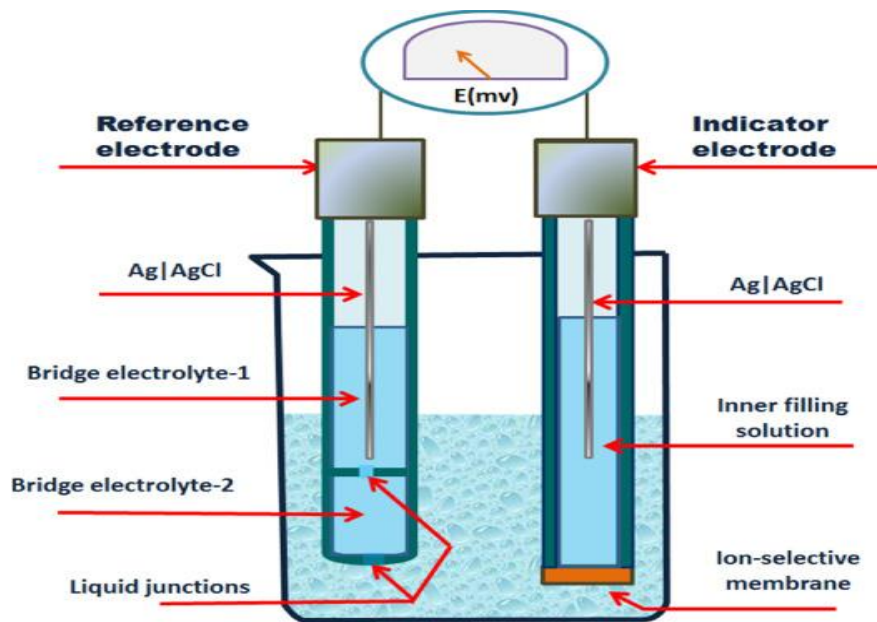


Figure (6): Diagram in schematic form. The typical potentiometric cell is depicted here using an ion-selective electrode in the accompanying (Abd El-Rahman & Salem, 2015).



Using Diazepam as a template, 2-acrylamido-2-methyl-1-propan sulfanic acid and N.N-methylene bis acrylamide as monomers and cross-linkers, respectively, as well as the benzoyl peroxide as an initiator, two electrodes were produced. Plasticisers are indispensable to ISE membranes. Membranes, including polymer and other materials compatibility ingredients, offer a membrane-homogeneous environment when plasticisers are utilised as membrane solvents. However, leaching of the plasticiser during The ISE membrane should see some practical use. be prevented because it would harm the electrode's performance over time. Because of a PVC matrix, four electrodes have been manufactured. The plasticisers Tritolyl phosphate (TTP) and dibutyl sebacate(DBS) are examples. The linearity range, correlation coefficients, detection limit (M), and life duration of the DZP-MIP1 (M1, M2) membrane-based electrodes were evaluated (day). The gathered information is presented in table 1 and figure 5.

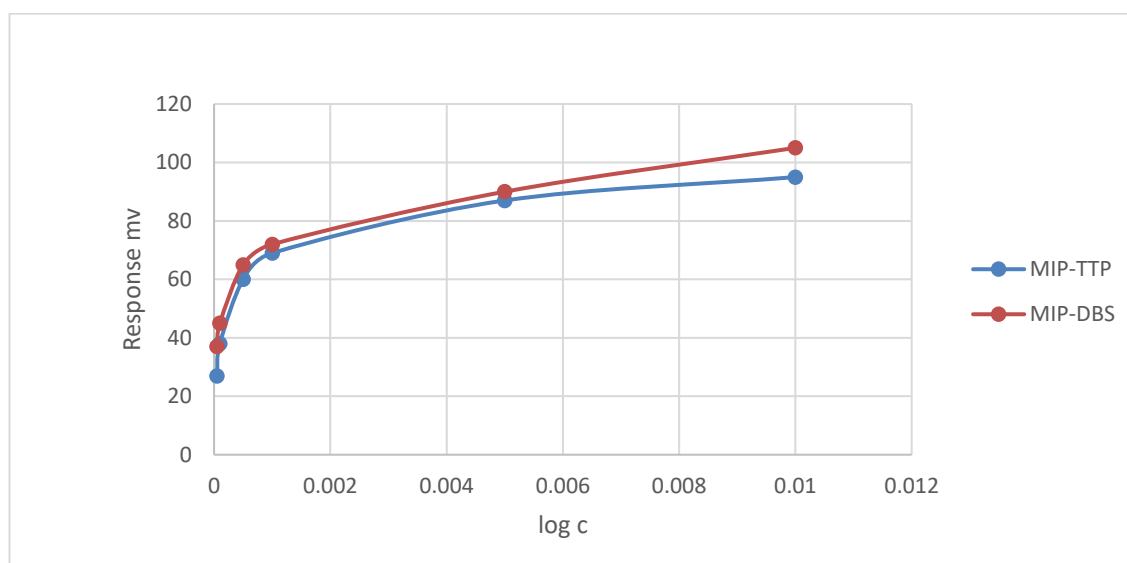


Figure (7): For DZP-MIP membrane electrodes, calibration curve.

Table (2): Characteristics of the diazepam-MIP electrode built with several functional plasticizers and monomers.

Membrane composition	DZP-MIP1+ TTP (M1)	DZP-MIP1+DBS (M2)
Slop (mV/decade)	27.113219	28.41902
Linearity range (M)	$5 \times 10^{-5} - 1 \times 10^{-2}$	$5 \times 10^{-5} - 1 \times 10^{-2}$
Correlation coefficient	0.9663	0.9932
The detection limit (M)	5×10^{-6}	5.5×10^{-6}
Lifetime (day)	19	16

Effect of pH on electrodes response:

Two electrodes were fabricated using Diazepam as a template and allyl chloride as a precursor. A pH investigation was conducted on DZP membrane electrodes with varying concentrations of Dia (5×10^{-3} and 5×10^{-4}). In pH investigations, pH testing (1-11) uses HCLacid(0.1M,1M) or NH4OH (0.1M,1M). As indicated in Table (3) and Figure 8, the end result achieved by including the proper amount of HCl/NH4OH the composition of electrodes is responsible for the difference in pH-value-related potential (Mahdi & Al-Bayati, 2020).

Table (3): Working pH range for Selective diazepam electrode.

Number and design of MIPs	Membranes	Membrane design	pH range	
			5×10^{-3}	5×10^{-4}
MIP DZP+ AMPS +N,N-MDAA	M1	DZP-MIP+TTP	3.5-7	5-7
	M2	DZP-MIP+DBS	3-4.5	2-5.5

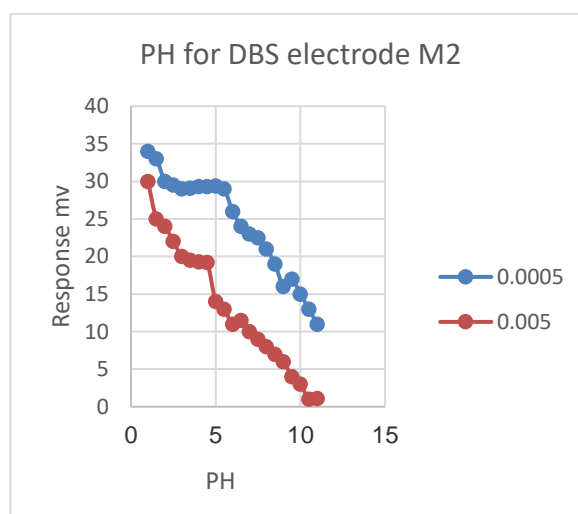
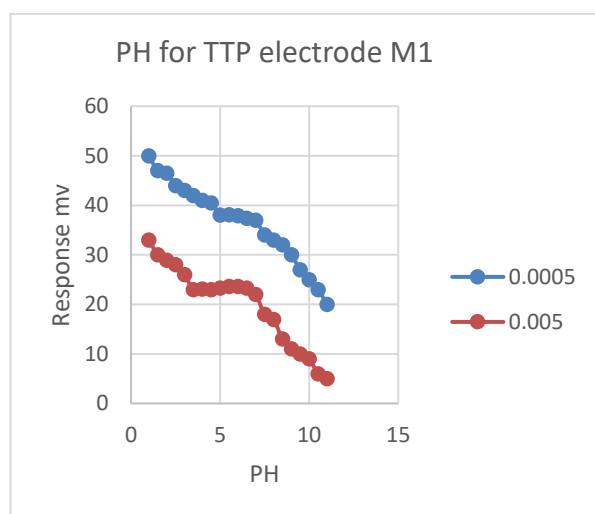


Figure (8): Effect of pH on the Diazepam [DZP-MIP+ TTP(M1) and DZP-MIP +DBS (M2)] electrodes at concentrations 5×10^{-3} and 5×10^{-4} .

Interference studies

Two electrodes were fabricated using Diazepam as a template and 2-acrylamido-2-methyl-1-sulfanic acid as a precursor. The pH For measuring the selectivity coefficient, a different solution approach was used. Utilise the unique equation required for these parameters, as shown in the following equation.

$$\text{Log } K_{\text{pot}} = \frac{(E_B - E_A)}{(2.303RT/zF)} + (1 - z_A/z_B) \log a_A \quad (\text{Al-Safi \& Al-Bayati, 2018})$$

EA, EB, zA, zB, and aA are, in order as well as the charges, potentials, and actions of the primary A ions and the interfering B ions when $a_A = a_B$.



The findings for principal ion selectivity coefficients and interference from other ions, such as (K^{+1} , Ca^{+2} and Al^{+3}), have been obtained in this investigation. Primary ion charge and secondary ion interference charge, as well as the concentration and composition of the electrodes, all have a role in determining the selectivity coefficients. All selectivity coefficient values were presented in Figures 9,10. Using several electrodes of the DZP membrane Tables 4,5 and

Table (4): Selectivity coefficients for (DZP-MIP+TTP) electrode at different concentrations of Diazepam.

Con.	Concentrations of Diazepam (M): concentrations of interference ions (M)					
	Interfering ions					
	K^{+1}		Ca^{+2}		Al^{+3}	
	E_B (mv)	$K_{A,B}$	E_B (mv)	$K_{A,B}$	E_B (mv)	$K_{A,B}$
10^{-2}	90	8.5169×10^{-2}	61.5	2.3934×10^{-3}	65.2	2.2325×10^{-3}
5×10^{-3}	88	1.0989×10^{-1}	60	1.0189×10^{-3}	63	6.1006×10^{-4}
1×10^{-3}	82	2.3601×10^{-1}	63	1.4862×10^{-3}	63	4.6987×10^{-4}
5×10^{-4}	75	2.7970×10^{-1}	65	1.1963×10^{-3}	51.6	8.2556×10^{-5}
1×10^{-4}	69	7.1196×10^{-1}	66	1.745×10^{-3}	52	7.7961×10^{-5}
5×10^{-5}	66	3.0167×10^{-1}	76	7.0533×10^{-3}	45.5	5.2863×10^{-5}

Table (5): Selectivity coefficients for (DZP-MIP+DBS) electrode at different concentrations of Diazepam.

Con.	Concentrations of Diazepam (M): concentrations of interference ions (M)					
	Interfering ions					
	K^{+1}		Ca^{+2}		Al^{+3}	
	E_B (mv)	$K_{A,B}$	E_B (mv)	$K_{A,B}$	E_B (mv)	$K_{A,B}$
10^{-2}	39	4.7600×10^{-3}	43	2.0814×10^{-3}	61	6.0959×10^{-3}
5×10^{-3}	33	9.8696×10^{-3}	39	1.6048×10^{-3}	51	1.9691×10^{-3}
1×10^{-3}	30	3.3275×10^{-2}	31	1.1410×10^{-3}	44	1.0343×10^{-3}
5×10^{-4}	27	4.6012×10^{-2}	24	3.6083×10^{-4}	32	1.4860×10^{-4}
1×10^{-4}	25	1.9781×10^{-1}	20	4.1716×10^{-4}	27	1.079×10^{-4}
5×10^{-5}	23	3.2164×10^{-1}	18	2.1450×10^{-4}	20	2.5212×10^{-5}

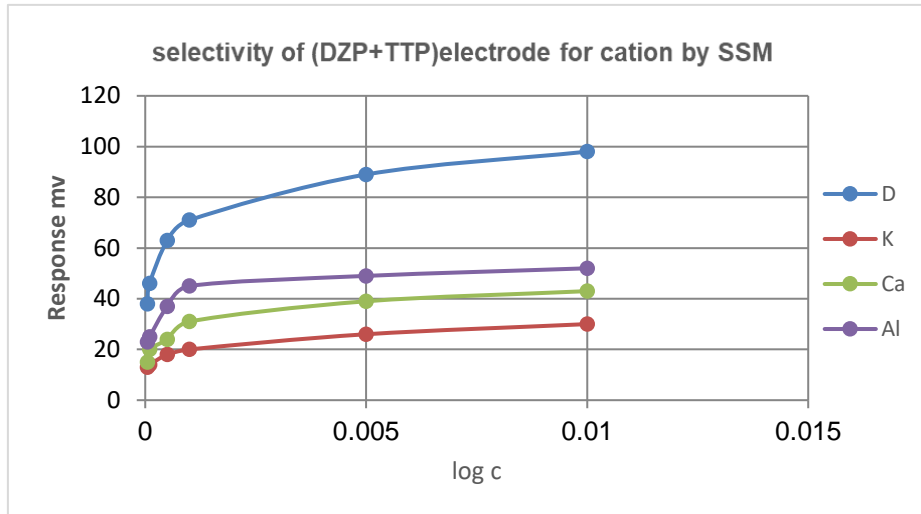


Figure (9): Selectivity of (DZP+TTP) ions at the electrodes using the Separation Solution Method.

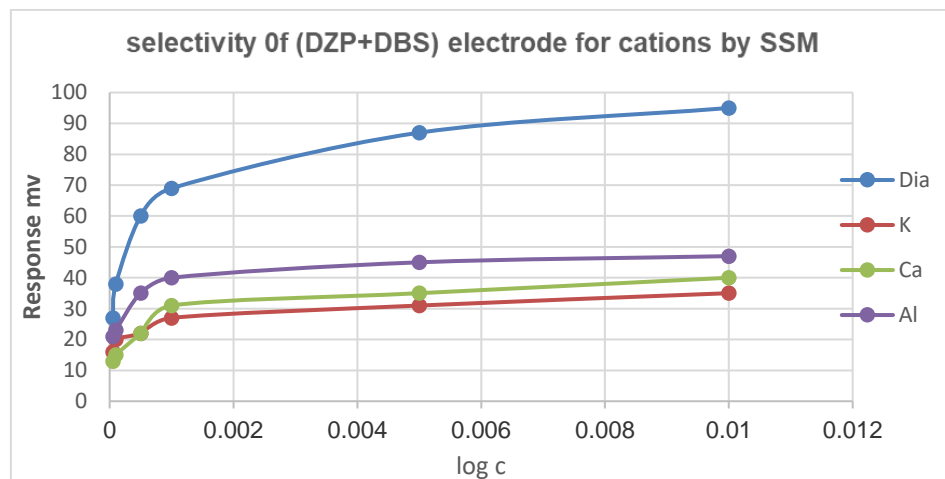


Figure (10): Selectivity of (DZP-DBS) ions at the electrodes using the Separation Solution Method.

Calculation by Multiple Standard Addition Method (MSA)

The concentrations employed in this method (5×10^{-3} and 5×10^{-4}) for graphing the antilog E/S (Y-axis) versus the normal level diazepam were investigated (X-axis). Figs. (9,10) depict what we found of diazepam ratios computed based on electrodes on DZP-MIP+ TTP, DZP-MIP+DBS.s performed on electrodes of DZP membrane utilising various concentrations of DZP.

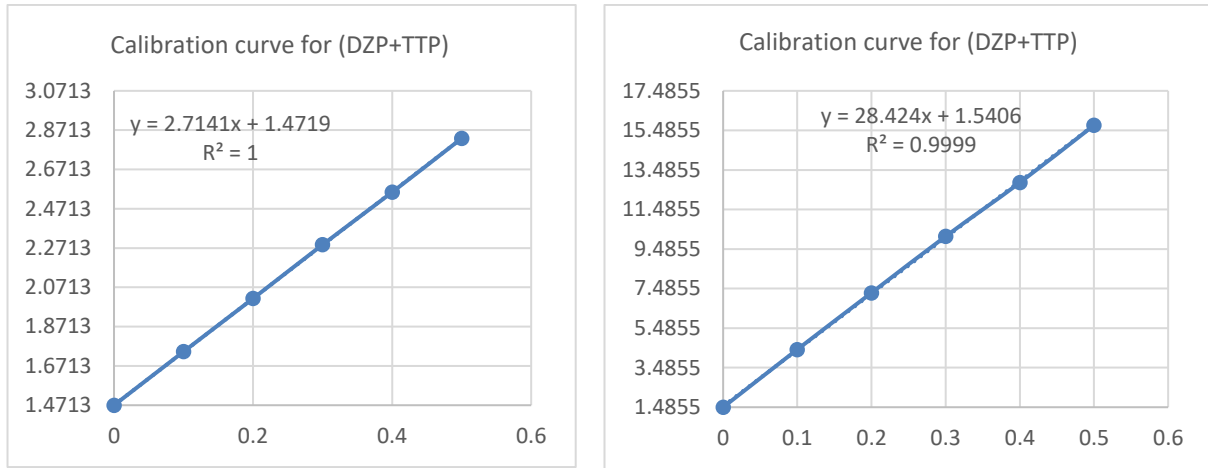


Figure (11): Antilog (E / S) in comparison to the total volume of the increased standard for the purpose of determining diazepam solution (5×10^{-3} and 5×10^{-4}) by MSA using (DZP–MIP + TTP) electrode.

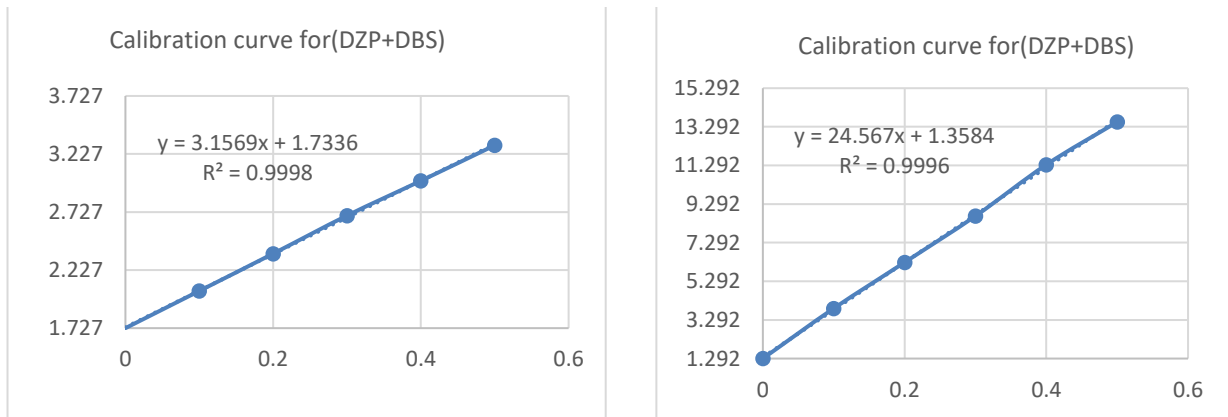


Figure (12): Antilog (E / S) in comparison to the total volume of the increased standard for the purpose of determining Diazepam solution (5×10^{-3} and 5×10^{-4}) by MSA using (DZP–MIP+ DBS) electrode.



Applications of pharmaceuticals.

Molecularly imprinted polymer-based ion selective electrodes were utilised to determine Diazepam in medicines. This ISE includes standard addition, direct, Gran plot, and multiple standard addition measurements. They were preparing solutions of Diazepam at 5×10^{-3} and 5×10^{-4} M concentrations. The RE%, RC%, and RSD% of Diazepam in medicinal use were calculated. The outcomes achieved are shown in Table (7).

Table (6): Determination of Diazepam Samples by Ion Selective electrodes (ISEs) techniques based on PVC membranes.

Electrode No.	Concentration (M)			
	Sample	Measurement using potentiometric methods		
		Direct	SAM	MSA
DZP-MIP+TTP	5×10^{-3}	5.0159×10^{-3}	4.9933×10^{-3}	4.9943×10^{-3}
	RSD%	0.85	2.89
	RC%	100.32	99.866	99.89
	RE%	0.32	-0.134	-0.11
	5×10^{-4}	5.0052×10^{-4}	4.9911×10^{-4}	4.9891×10^{-4}
	RSD%	0.4845	0.37
	RC%	100.40	99.822	99.78
	RE%	0.10	-0.178	-0.22

Electrode No.	Concentration (M)			
	Sample	Measurement using potentiometric methods		
		Direct	SAM	MSA
DZP-MIP+DBS	5×10^{-3}	5.0250×10^{-3}	4.9881×10^{-3}	4.9893×10^{-3}
	RSD%	0.82	2.76
	RC%	100.50	99.762	99.79
	RE%	0.50	-2.38	-0.21
	5×10^{-4}	4.9871×10^{-4}	4.9856×10^{-4}	5.0092×10^{-4}
	RSD%	0.911	0.52
	RC%	97.94	99.712	100.18
	RE%	-2.06	-0.288	0.18

Table 7: Sample analyses of pharmaceutical Diazepam using DZP-MIP+TPH electrode.

pharmaceutical	(Iraq)		
	Direct	SAM	MSA
Concentration prepared	5×10^{-3}	5×10^{-3}	5×10^{-3}
found	4.9176×10^{-3}	4.9933×10^{-3}	4.9649×10^{-3}
RC%	98.35	99.867	99.30
RSD%	1.2674	2.89
RE%	-1.65	-0.134	-0.70
pharmaceutical	Direct method	SAM	MAS
Concentration prepared	5×10^{-4}	5×10^{-4}	5×10^{-4}
Found	5.0198×10^{-4}	5.0341×10^{-4}	4.9624×10^{-4}
RSD%	1.6953	1.47
RC%	100.40	100.682	-0.75
RE%	0.40	0.628	99.25

pharmaceutical	(Iraq)		
Concentration prepared	Direct	SAM	MSA
	5×10^{-3}	5×10^{-3}	5×10^{-3}
found	4.9577×10^{-3}	4.9881×10^{-3}	4.9855×10^{-3}
RC%	99.15	99.762	99.71
RSD%	1.1893	2.76
RE%	-0.85	-0.238	-0.29
pharmaceutical	Direct method	SAM	MAS
Concentration prepared	5×10^{-4}	5×10^{-4}	5×10^{-4}
Found	4.818×10^{-4}	4.9838×10^{-4}	5.0473×10^{-4}
RSD%	2.249	3.6907
RC%	96.36	99.676	100.95
RE%	-3.64	-0.324	0.95

*each measurement was carried out three times.

CONCLUSION

By combining different plasticisers with Diazepam membranes, selective electrodes can be created. TTP and DBS plasticisers were employed to manufacture PVC-based diazepam membrane electrodes. The results obtained for all electrodes applied to standard and medicinal solutions were excellent. Developing electrodes for use in pharmaceutical analytical determination is intended to serve this purpose.

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