

Synthesis and Characterization of N-(2-acetylthiophene) salicylideneimine (ATS) as Ionophore for Polymeric Membrane Ag (I) Selective Electrode

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A NOVEL silver ion selective membrane electrode which was based on a newly synthesized N-(2-acetylthiophene) salicylideneimine (ATS) by the reaction of 2-acetyl thiophene with 2-aminophenol at 60°C in ethanol by using ultrasound irradiation. ATS was characterized by elemental analyses, FTIR, MS and ¹H-NMR. The best performance was obtained for the membrane of composition (ATS): PVC: DOS in the ratio 1:33:66 wt %. The sensor shows a linear potential response for Ag⁺ over a wide concentration range 6.0 x 10⁻⁸ -1.0 x 10⁻¹ M and detection limit 4.36 x 10⁻⁸ with Nernstian slope (57.93 mV decade⁻¹ of activity) between a wide pH range (3.0-10) and a fast response time of 10s. The selectivity coefficient values were calculated with separate solution method, which indicate good selectivity of Ag⁺ ion. The sensor has a lifetime of five weeks and could be used as an indicator electrode in the potentiometric titration of Ag⁺ vs. stander solution of NaCl and in the estimation of Ag⁺ in photographic solutions and medical X-ray samples.

Keywords: Poly vinyl chloride (PVC), N-(2-acetylthiophene) salicylideneimine (ATS), Silver selective sensor and Nernstian slope.

Silver (I) ion has been used in a wide range of products such as electronic products, alloys, jewelry and photographic film. Also due to its antibacterial effect, accelerating wound healing and anti-inflammatory properties, silver and its compounds have been applied in the medical and health care field ⁽¹⁾. Silver salts have also been used to disinfect water used for drinking and recreational purposes, in dental and pharmaceutical antibacterial ⁽²⁾. However, it is also well known that silver inactivates sulfhydryl enzymes, combines with amine, imidazole and carboxyl groups of various metabolites. It has been found that an excess of silver is toxic to fish and microorganisms at a concentration as low as 0.17µg/l ⁽³⁾. Thus, the increasing use of silver compounds in industry and medicine has resulted in an increased silver content of environmental samples.

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On the other hand, superfluous silver and its compounds have toxic effect on both human and animal health. Consequently, determination of the trace amount of silver ions in samples is necessary. Several techniques such as atomic absorption spectroscopy^(4,5) inductively coupled plasma mass spectroscopy⁽⁶⁻⁹⁾ and flow injection analysis^(10,11) are used for its determination at low concentration level. There has been a growing interest in the development and application of potentiometric sensors for monitoring concentration of various species in biological, environmental, and industrial fields. Because of their unique advantages such as fast response, ease of preparation, low cost, possibility of using in complex and color media, wide linear dynamic range, relatively low detection limit, and adequate selectivity for most analyses⁽¹²⁾. Thus, a number of Ag(I) sensor based on crown ethers⁽¹³⁻¹⁵⁾ calix[4]arene derivatives⁽¹⁶⁻¹⁸⁾, bis-pyridinetetramidemacrocycle⁽¹⁹⁾ bis(dialkyldithiophosphate) derivative⁽²⁰⁾ and bis(thiothiazole)derivative⁽²¹⁾ Schiff bases^(22, 23) in PVC have been reported. It is well known that the soft heavy metal ions, Ag (I), Pb (II) and Hg (II) display great affinity for soft coordination centers like sulfur and nitrogen. By using ionophores containing sulfur and nitrogen atoms in ion selective electrodes, the electrode are sensitive to soft heavy metal ions and the selectivity for soft heavy metal ions against alkali metal ions is significantly increased⁽²⁴⁾. In this study, we utilized N-(2-acetylthiophene) salicylideneimine (ATS) ionophore, as Schiff base selective carrier for silver ions. The results obtained with (ATS) based on silver electrodes showed high selectivity coefficients for silver ion over all the studied alkali, alkaline, transition ions and heavy metal ions.

Experimental

Equipments

Elemental microanalyses of the separated Schiff base for C, H, N and S were performed in Main Defense Chemical Laboratory, Almaza, Cairo, Egypt. The FTIR spectra (400 – 4000 cm^{-1}) of the compound was recorded as KBr discs using a Shimadzu 8000 FTIR spectrometer at the central laboratory, Ain Shams University, Cairo, Egypt. ¹H-NMR spectra recorded using a Varian spectrometer, 300 MHz in Main Defense Chemical Laboratory. All potentiometric measurements were performed at room temperature using pH/mV meter [Orion model A 720], double junction Ag/AgCl reference electrode [Orion model 90-02] containing 10% KNO_3 in the outer compartment. Combination glass electrode [Orion model 8-02] was used for all pH measurements.

Ag/AgCl / internal solution (1mM NaCl in acetate buffer pH=4.0)/ PVC membrane / test solutions / AgCl, Ag

Reagents and materials

2-acetylthiophene was obtained from Merck, and 2-aminophenol was obtained from Aldrich (USA). Solvents and reagents were purified and dried by distillation. High molecular weight poly vinyl chloride (PVC), 2-ethyl sebacate (DOS), potassium tetrakis (4-chlorophenyl) borate (KTpClPB) and

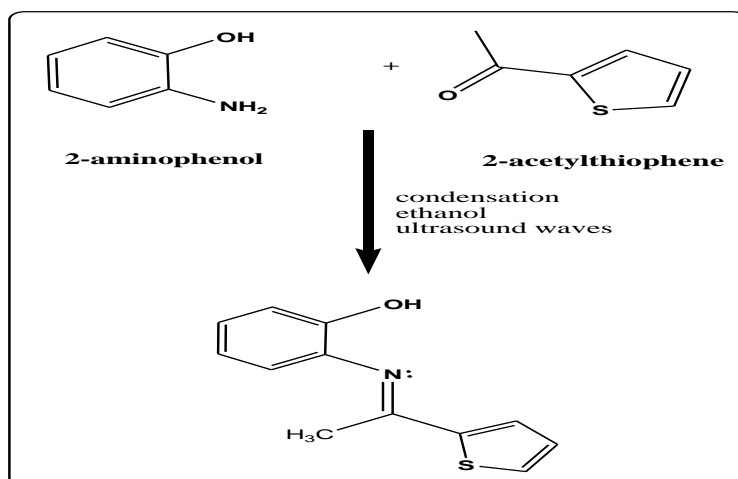
tetrahydrofuran (THF), o-nitrophenyloctylether (O-NPOE), dioctylphthalate (DOP) were obtained fluke (Ronkonkoma, NY). All other reagents used as purchased without further purification. Deionized water was used in the preparation of buffer solutions. 0.1 M stock solutions were prepared by dissolving AR grade metal nitrates solution in buffer solutions and standardized wherever necessary. The working solutions of different concentration were prepared by stock solution.

Syntheses of N-(2-acetylthiophene)salicylideneimine Schiff base (ATS)

The ligand was synthesized by slow addition of 2-acetylthiophene (0.33 ml, 3 mmole) in 30 ml ethanol to 2-aminophenol (0.327 gm, 3 mmole) in 30 ml ethanol. The reaction mixture was heated to reflux at 60°C and subjected to ultrasound irradiation to for 2hr. The brown product obtained after cooling was filtered off and washed with few amounts of ethanol then diethylether. Fine crystals were obtained by recrystallization from hot ethanol. The yield was (0.526gm, 74.60 %), m.p. (157-159°C).

Anal. Calcd. For C₁₂H₁₁NSO: C, 66.36; H, 5.07; N, 6.45; S, 14.75. Found: C, 66.40; H, 4.87; N, 6.60; S, 14.60%. Infrared spectrum (cm⁻¹ KBr disk); 3304 (v.s) (ν_{OH}, phenolic), 3054(s)(ν_{C-H}, Ar.); 2958, 2851 (v.s)(ν_{C-H} asym. & sym., Aliph.); 1602 (s) (ν_{C=N}, azomethine); 1141 (s) (ν_{C-O}, phenolic); 1467(m) (ν_{C=C}, Ar.); 743(v.s) (ν_{C-S}, thiophenolic).

(¹H-NMR, DMSO-d₆, ppm); δ 3.172 (s, 3H, CH₃), 6.358 – 6.986 (m, 4H, Ar-H), 7.365 – 7.722 (m, 3H, thiophene protons), 8.879 (s, 1H, OH). MS (m/z, %); 217 (M⁺, 70.4); 57 (100), 69 (65.17), 76 (12.36), 82 (12.36), 94 (60.67), 107 (76.40), 134 (61.80), 175 (64.04), 175 (64.04), 187 (60.67) and 202 (65.17). The reaction for the formation of (ATS) is illustrated in Scheme 1.



Scheme 1. Schematic representation for the formation of Schiff base ligand, (ATS).
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Membrane preparation

The PVC based membranes were prepared as described elsewhere⁽²⁵⁾, by dissolving appropriate amounts of Schiff base (TAS), anion excluder KTpClPB, solvent mediators 2-nitrophenyloctylether (NPOE), dioctyl phthalate (DOP), 2-ethyl sebacate (DOS) and appropriate amount of PVC in THF (2mL). After complete dissolution of all the components and thorough mixing, the resultant cocktail was transferred to a glass ring placed on a glass plate. Membranes were formed after controlled evaporation of THF at room temperature overnight. A flexible, transparent membrane of a thickness of ~ 0.2–0.4 mm was obtained. The discs of 7 mm diameter were cut using a cork borer, and then were glued to a PVC tip which was clipped to the end of the electrode body. The PVC membrane electrodes were pre-conditioned by immersion in a 1 mM NaCl for at least 24 hr prior to use. Membranes of different compositions were prepared and investigated. Those, which gave reproducible results and best performance characteristics, were selected for detailed studies. The optimum compositions of best performing membranes are given in (Table 1).

Sample pretreatment

The waste photographic and medical x-Ray films solutions were prepared for the measurement of their silver content as follows. 20 ml of the sample was added to 10 ml of 3 M nitric acid and 40 ml distilled water. The solution was boiled for 1 hr. min until its volume was reduced to half of its original values. The resulting solution was neutralized with sodium hydroxide solution to a pH 4.0, and filtered. The filtrate and washings were diluted to 100 ml in a volumetric flask with acetate buffer pH 4.0 and used for potentiometric and atomic absorption measurements. A portion of the film sample 2.0 g was mixed with 5 ml of concentrated HNO₃ and 50 ml water and boiled for ~1 h. The resulting mixture was then filtered and the filtrate was neutralized with 1 M sodium hydroxide solution and diluted to 100 ml using acetate buffer pH 4.0.

Results and Discussion

Influence of membrane composition on the potentiometric response

The influence of addition of various mol % anionic sites (relative to ATS ionophore) and using different types of plasticizers (o-NPOE, DOP and DOS) were investigated. Surprisingly we found that all membranes (E₁, E₂) formulated with high polarity plasticizers o-NPOE and DOP and (E₄, E₅) formulated with DOS (with anionic additives) exhibited minimal cationic response to all cations investigated including primary ion (silver). As depicted in (Table 1 & Fig. 1), membrane sensor plasticized with low polarity DOS (E₃ without additives) showed better response slopes. This is consistent with literature finding that polymeric membrane of low-polarity exhibited improved responses toward monovalent cations⁽²⁶⁾.

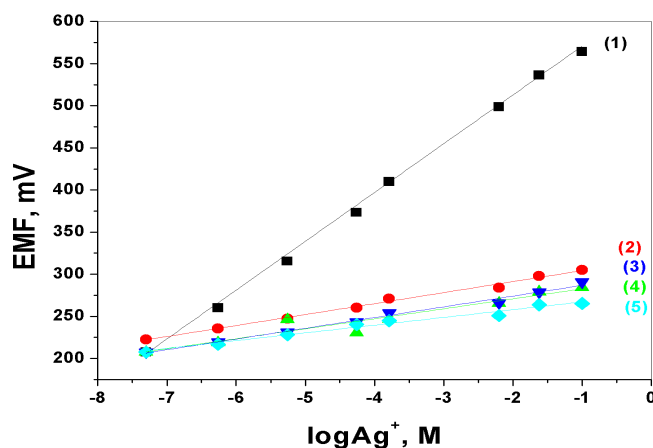


Fig. 1. Potentiometric response, measured in 50mM acetate buffer pH 4.0, of various compositions of ATS-based membrane electrode toward silver ions: E₃(1), E₄(2), E₅(3), E₁(4), E₂(5).

TABLE 1. Effect of the plasticizer type and addition of ionic sites on the response characteristics of silver membrane electrodes based on ATS ionophore.

Electrode No.	Membrane Composition					Slope mV/decade	Linear Range M	Detection limit M	R ²
	PU mg	PVC mg	Plasticizer (132 mg)	ATS mg	(TCIPB) mol%				
E ₁	-	66	o-NPOE	2	12.75	6.0x10 ⁻⁸ -1.0x10 ⁻¹	4.89x10 ⁻⁸	0.991
E ₂	-	66	DOP	2	9.16	6.0x10 ⁻⁸ -1.0x10 ⁻¹	5.13x10 ⁻⁸	0.983
E ₃	-	66	DOS	2	57.93	6.0x10 ⁻⁸ -1.0x10 ⁻¹	4.36x10 ⁻⁸	0.992
E ₄	-	66	DOS	2	10	13.03	1.0x10 ⁻⁷ -1.0x10 ⁻¹	5.13x10 ⁻⁸	0.981
E ₅	-	66	DOS	2	50	12.54	5.0x10 ⁻⁸ -1.0x10 ⁻¹	4.47x10 ⁻⁸	0.958

R² : Correlation coefficient

Selectivity coefficients of optimized silver ion-sensor

The potentiometric selectivity of the optimized sensor was investigated for several ions and the selectivity coefficients were calculated using separated sample solution method (SSM)⁽¹²⁾. The cationic response of the optimized potentiometric sensor (E₃) towards various cations is shown in Fig. 2, and the selectivity coefficient is shown in Table 2. As can be seen optimized potentiometric sensor (E₃) exhibited high selectivity for Ag⁺ ion over a number of monovalent, divalent and trivalent cations, also the optimal membrane sensor showed remarkably better selectivity in comparison with membrane sensor prepared with anionic sites (E₄ and E₅). This result agreed very well with the view that the soft coordination site of sulfur offers great affinity towards silver which is soft (d¹⁰) transition metal ion⁽²⁷⁾.

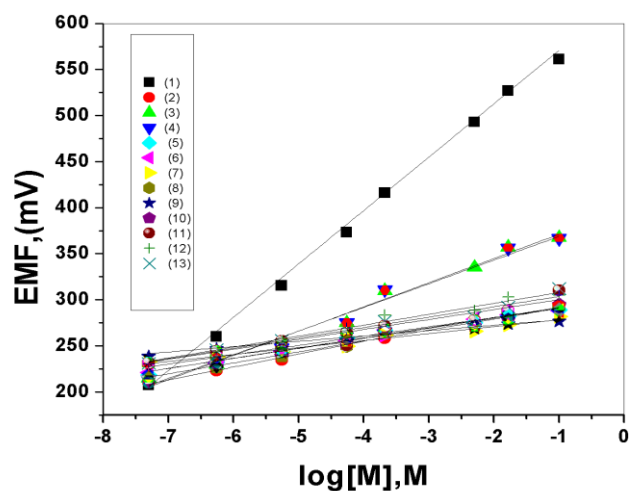


Fig. 2. Potentiometric response of ATS-based silver-selective membrane electrode E_3 , measured in 50 mM acetate buffer pH 4.0 towards various metal ions : Ag^+ (1), Li^+ (2), Na^+ (3), Hg^{2+} (4), Cu^{2+} (5), Pb^{2+} (6), Mn^{2+} (7), Zn^{2+} (8), Ni^{2+} (9), Co^{2+} (10), Al^{3+} (11), Sr^{3+} (12), Cr^{3+} (13).

TABLE 2. Selectivity coefficients $\log K_{Ag^+, X^{n+}}^{pot}$ for optimized silver-membrane sensor E_3 measured in 0.05 M acetate buffer, pH 4.0.

Metal ion	$\log K_{Ag^+, X^{n+}}^{pot}$
Li^+	-4.4
Na^+	-4.8
Hg^{2+}	-4.9
Cu^{2+}	-5.2
Pb^{2+}	-5.2
Mn^{2+}	-5.2
Zn^{2+}	-5.2
Ni^{2+}	-5.2
Co^{2+}	-5.5
Al^{3+}	-5.6
Sr^{3+}	-5.7
Cr^{3+}	-5.9

Effect of pH

The pH dependence of the optimized membrane sensor (E_3) was investigated at two concentrations of Ag^+ (1.0×10^{-3} and 1.0×10^{-4} M) solution over a pH range of 2 to 11 and the results obtained are given in Fig. 3. It is seen from this figure that the potential remains constant over a pH range of 3 to 10. This means that the optimized sensor can be used to measure a wide range of environmental and industrial water samples without pH adjustment.

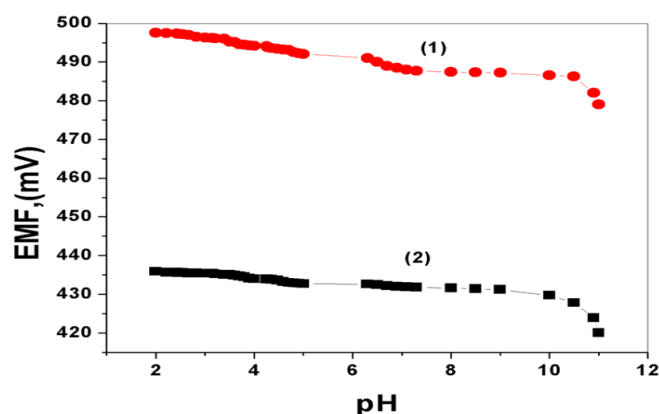


Fig. 3. Effect of pH on the potentiometric response of optimized membrane electrode E_3 measured in 50 mM acetate buffer pH 4.0, in the presence of 1.0×10^{-3} M (1) and 1.0×10^{-4} M (2) of AgNO_3 .

Response mechanism and lifetime

Figure 4, shows the response time of the optimized sensor (E_3) when exposed to increasing silver ion concentrations from $1.0 \mu\text{M}$ to 0.1 M in 50 mM acetate buffer, pH 4.0. As can be seen, over the investigated concentration range, the optimized sensors reached to the equilibrium response in a very short time ($<10\text{s}$). The life time of the optimized membrane electrodes was found to have the same slope with deviation of $\pm 4 \text{ mV/decade}$ over a period of five week.

Potentiometric titration

The optimized selective membrane sensor (E_3) was utilized as an indicator electrode in the potentiometric titration of different concentrations of silver ions with standard sodium chloride solution. Typical titration curves with sharp inflection breaks were obtained as can be seen in Fig. 5. The signal change at the inflection breaks was found to be in the range 390-470 mV depending on the starting silver ion concentration.

Analytical application

The optimized of the ATS-based silver electrode was further assessed by measuring the silver ion concentrations in acetate buffered, pH = 4 photographic solution samples and medical x-Ray films. Comparison of these results with the data obtained using atomic absorption spectrometry (AAS) showed very good agreement as shown in Table 3.

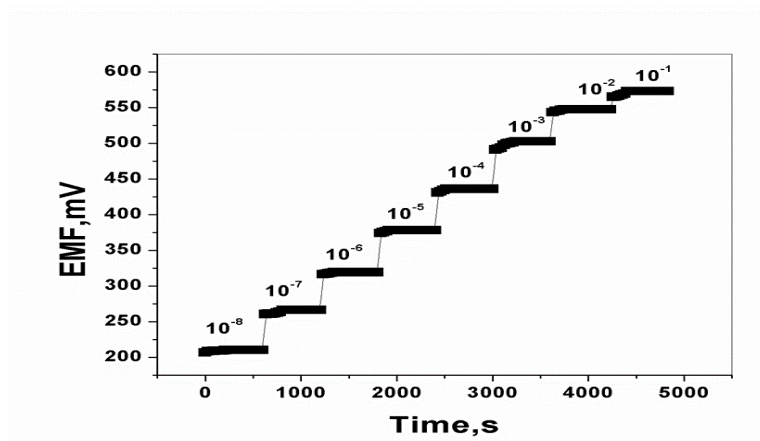


Fig. 4. Typical response time of optimized electrode E₃ measured in 50 mM acetate buffer, pH 4.0 at different silver concentrations

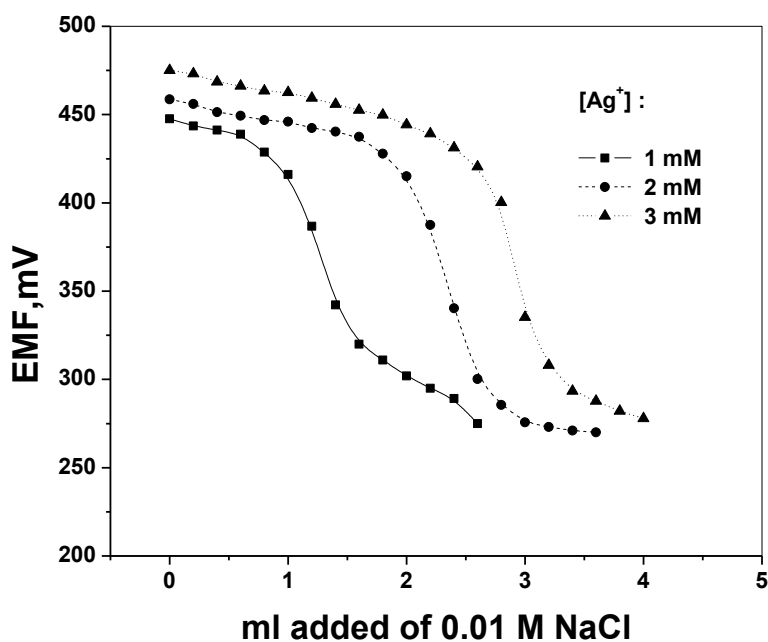


Fig. 5. Typical potentiometric titrations of different amount of silver ions with standard sodium chloride solution monitored using optimized membrane electrode E₃.

TABLE 3. Determination of Ag⁺ ions in photographic and x-rays samples using optimized sensor (E₃) in comparison with standard atomic absorption method.

Real samples	AAS±SD ^a mg/l	Potentiometry±SD ^a mg/l	Recovery %
Fixing solutions			
A	11.9 ± 0.4	11.6 ± 0.4	97.47
B	6.3± 0.9	6.17 ± 1	97.94
Medical X-rays ^b			
A	2.06 ± 0.3	2.03 ± 0.4	98.54
B	3.47 ± 0.4	3.47 ± 0.5	100.00

a : Mean value ± standard deviation (SD) {two determinations }.

b: 2 mg samples were dissolved in 10 ml solution.

Conclusion

The investigations demonstrate the utility of the membrane sensor incorporating N-(2-acetylthiophene)salicylideneimine (ATS) as ionophore in determining Ag⁺ in the concentration range 6.0×10^{-8} - 1.0×10^{-1} M. The functional pH range is 3.0 to 10. It could be used as an indicator electrode in the potentiometric titration of Ag⁺ ions with NaCl solution. Analytical application shows that it holds promise for regular and routine analysis of Ag⁺ in diverse samples. It exhibits good reproducibility, fast response time and can be used for five week.

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تحضير وتوصيف [٢-اسيتيل ثيوفين) سالسليدين أمين] كمجس غشائى لتقدير الجهد الكهربى لعنصر الفضة

ايمان حامد عبد الناصر، مناره احمد أيوب، منى عبد العزيز أحمد و
مريم جرجس رزق
قسم الكيمياء- كلية البنات للآداب والعلوم والتربيه- جامعه عين
شمس - القاهره - مصر.

تحضير مجس غشائى جديد لتقدير الجهد الكهربى لعنصر الفضة . وهذا المجس
يحتوى على [٢-اسيتيل ثيوفين) سالسليدين أمين] وقد تم تحضيره بتكثيف ٢-اسيتيل
ثيوفين مع ٢-امينو فينول وقد تم اثبات تحضير المجس باستخدام التحليل العنصرى
و طيف إمتصاص الأشعه تحت الحمراء و طيف الكتله و طيف الرنين النووى
المغناطيسى ونكوين افضل مجس من نوع عديد (كلوريد الفينيل) و ثنائى أثيل
أوكتيل سيباكات كماده ملدنه. و يظهر هذا المجس أستجابته سريعه لمحاليل الفضة و
يعطى أستجابته سريعه بميل كاتيونى نرنستى قدره ٥٧.٩٣ مللى فولت لكل تركيز
عقدى فى مدى واسع للأس الهيدروجينى من (٣ إلى ١٠) و قد أمكن أستخدام
هذا المجس فى تقدير أيون الفضة فى المحاليل الفوتوغرافيه و صور الأشعه السينيه
الطبيه. و قد تم الحصول على نتائج جيده مطابقه للنتائج التى تم الحصول عليها
بأستخدام قياس طيف الأمتصاص الذرى لأيون الفضة كطريقه قياسيه.