



Cite this: *Analyst*, 2024, **149**, 5225

A sulphide resistant Ag|AgCl reference electrode for long-term monitoring†

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Reference electrodes which demonstrate long-term potential stability are essential for many continuous monitoring applications and are commonly based on Ag|AgCl electrodes; however, these electrodes are susceptible to poisoning from aqueous sulphide species which are commonly present in wastewater and natural groundwater. This work presents a sulphide resistant solid-state reference electrode (SSRE) based on a composite material using suspended KCl electrolyte and sacrificial AgCl in a cross-linked polyvinyl acetate polymer matrix. Sulphidation of the sacrificial AgCl produces a stable Ag₂S precipitate and prevents further ingress of the poisoning sulphide species through the composite material. A novel SSRE using this material is compared to a control SSRE without suspended AgCl and a typical liquid filled reference electrode. These three reference electrodes are studied using electrochemical impedance spectroscopy (EIS), and their application is also studied in potentiometric pH sensing and cyclic voltammetry (CV). The long-term sulphide resistance of the two SSREs is also studied with potentiometry, and cross-sections of these electrodes were examined using micro X-ray fluorescence (μXRF). Both SSREs demonstrated higher impedance than the liquid reference electrode but were similar to other SSREs reported in the literature. This impedance did not result a meaningful difference in potentiometric pH sensing or CV experiments done using typical scan rates. The KCl/AgCl SSRE exhibited remarkable sulphide resistance, with all samples demonstrating a stable potential without maintenance after ca. 120 days of continuous immersion in 1 g L⁻¹ Na₂S solution, whereas KCl SSRE samples all demonstrated significant drift before this time. μXRF sulphur maps revealed that suspended AgCl prevented sulphide ingress, thus protecting the embedded Ag|AgCl electrode. This work presents a reference electrode that could enable long-term monitoring in challenging sulphide solutions, and also highlights a novel approach for preventing reference electrode poisoning which could be more widely explored.

Received 9th August 2024,
 Accepted 12th September 2024

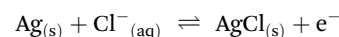
DOI: 10.1039/d4an01076k

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Introduction

Reference electrodes are needed in most electroanalytic applications, ideally proving a stable reference for the measurement of indicator or working electrode potentials. There are numerous mercury and silver based reference electrodes for aqueous systems, but Ag|AgCl reference electrodes are most commonly used as the materials are inexpensive and relatively safe.¹ The potential of these reference electrodes is dependent on the chloride activity at the internal Ag|AgCl electrode as described

by the Nernst equation,² and has the following half-cell reaction:



Drift in reference electrode potential is therefore often caused by chloride electrolyte leakage, however another source of instability is contamination of the Ag|AgCl electrode by chemical interferants such as hydroxide or sulphide species.³

Typical liquid-filled reference electrodes provide adequate performance in most laboratory applications, however there is an ongoing need for new designs which can withstand harsher chemical and physical conditions. In particular, *in situ* environmental monitoring relies on having sensors which require minimal maintenance or calibration,^{4–7} otherwise their use becomes less attractive compared to laborious sampling and laboratory analysis methods. For electrochemical sensors the reference electrode is often a limiting factor in online monitoring,⁸ especially for potentiometric

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†Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4an01076k>



sensors where the indicator electrode potential is directly used to determine analyte concentration.

Solid state reference electrodes (SSREs), constructed without an internal liquid reservoir, have proved to be an effective approach to making a more robust reference electrode. Since the earliest reports of SSREs in the late 1990s,^{9,10} numerous examples have since been reported in the literature including many polymer/electrolyte composite reference electrodes,^{11–13} and printed reference electrodes.^{14–17} Other recent reference electrode designs include a self-referencing pulstrode which relies on periodically reducing a AgI coating and using the resulting electromotive force as a reference,^{18,19} and capillary based electrodes which prevent contamination by using low flow rates and small sample volumes.^{20,21} It is clear from the wide array of designs that it is perhaps impossible to create an ideal reference electrode suitable for every condition, and therefore specifically tailored options are needed for each use case.

Sulphide species are often found in wastewater and natural groundwater,^{22–24} making poisoning an ongoing challenge for electrochemical sensors in these environments. Sulphidation of AgCl to form Ag₂S has been widely reported in the literature,^{25–28} with the later material demonstrating a low solubility product and therefore great stability.²⁹ Previously we reported a SSRE which demonstrated a continuous working lifetime of 175 days without maintenance in 1 M Na₂SO₄ solution.³⁰ In this work we build on this design exploiting the creation of the aforementioned stable metal sulphide to create a sulphide resistant reference electrode. Sacrificial AgCl is suspended with KCl electrolyte and results in the precipitation of aqueous sulphide species as Ag₂S before they can diffuse through the polymer matrix and contaminate the Ag|AgCl electrode (see Fig. 1). This simple design can be used similarly to other existing reference electrodes without a need for significant changes in electronics or incorporation of liquid sampling equipment. Electrochemical impedance spectroscopy (EIS) is used to study the KCl/AgCl SSRE, with comparisons to our previous KCl SSRE and a commercial liquid filled reference electrode. These three reference electrodes are

also compared in two applications: potentiometric sensing with a glass pH electrode, and cyclic voltammetry (CV) experiments using [Ru(NH₃)₆]Cl₃. Finally, the long-term sulphide resistance of both SSREs is studied using potentiometry and micro X-ray fluorescence (μXRF).

Experimental

Materials

Silver wire (99.9%) was purchased from A&E Metals. Vinyl acetate (99%), ethylene glycol dimethacrylate (EGDMA) (98%), and Na₂S (98%) were supplied by Aldrich. KCl (99.8%) and glass chromatography vials were supplied by Thermo Fischer Scientific. AgCl (99.9%) was supplied by Alfa Aesar. 2,2'-Azobis (2-methylpropionitrile) (AIBN) (98%) was supplied by Acros Organics. pH buffers were supplied by Merck. [Ru(NH₃)₆]Cl₃ (99%) was supplied by Strem Chemicals. Insulated copper wires, crimps, and heat shrink tubing were purchased from RS Components. The liquid reference electrode used for comparisons in EIS, potentiometric, and CV experiments was a saturated KCl Ag|AgCl (1004 Series, Koslow).

Electrode preparation

Ag|AgCl electrode preparation. A silver wire (4 cm × 2 mm) was first electrochemically coated with AgCl in a 2-electrode cell containing an aqueous 1 M KCl solution and a platinum flag counter electrode (10 mm × 9 mm). The silver wire was oxidised using constant current with a current density of 3.65 mA cm⁻² for 100 s, creating a dark grey coating. Scanning electron microscopy images of wire cross-sections display a AgCl coating thickness of approximately 2 μm (see Fig. S1a†).

Polymerisation and electrode preparation. Monomer solutions were prepared by dissolving AIBN (100 mg) in vinyl acetate (9.8 mL) with EGDMA (200 μL). KCl SSRE slurries were prepared by suspending KCl (11.33 g) in the monomer stock solution. KCl/AgCl SSRE slurries were similarly prepared by suspending KCl (11.33 g) and AgCl (500 mg) in the monomer stock solution. Each electrode had 1.5 mL of this slurry pipetted into a 1.5 mL glass chromatography vial (32 mm × 11.6 mm). The rubber septum of the vial's lid was pierced with the AgCl-coated Ag wire then screwed tightly onto the vial, such that the coated wire was immersed in the slurry and sat 5 mm from the bottom of the vial. The vial was then placed in a convection oven set to 50° C and left overnight to initiate the polymerisation. KCl SSREs presented a solid white composite, and KCl/AgCl SSREs were a slightly darker grey solid composite. SEM images of the cured polymer composites demonstrated that KCl and AgCl particle sizes were all <500 μm (see Fig. S1b†). The bottom of the vial was ground away using a ceramic tile saw to expose a hard composite material, leaving the submerged end of the wire approximately 3 mm from the opening. Insulated copper wires were then crimped onto the exposed silver contact, then the connection was insulated using 3 mm diameter heat shrink tubing and embedded in epoxy resin. Excluding the long-term sulphide resistance tests

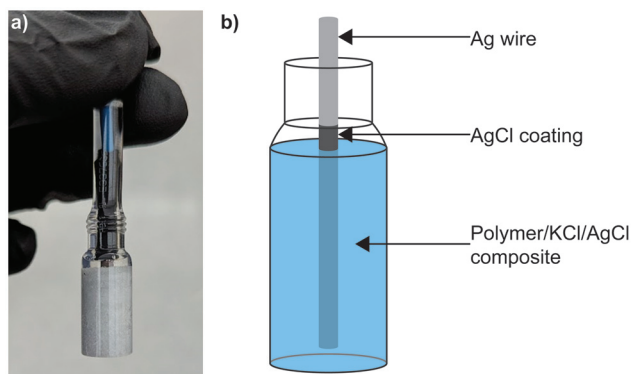


Fig. 1 (a) picture of KCl/AgCl SSRE, and (b) schematic illustrating the electrode's construction.



where SSREs were conditioned *in situ*, SSREs were soaked in deionised water for one week then kept in 3 M KCl solution prior to electrochemical testing. After this conditioning the polymer composite slightly softened but remained firm and inelastic for both SSRE types.

Physical characterisation

Scanning electron microscopy (SEM). Imaging of unconditioned SSRE cross-sections was done using a FEI Quanta 400, field emission, environmental scanning electron microscope (ESEM). Initially, samples were sectioned, mounted in Struers Epofix 2-part epoxy resin and mechanically polished using traditional metallographic techniques down to a final grit size of 1 μm . Samples were then coated with a 10 nm layer of carbon to dissipate charge, using an Edwards Auto 306 carbon arc, vacuum coater. After insertion into the ESEM and using high vacuum imaging mode, backscattered electron images of the sample were collected at a beam energy of 10 keV and a probe current of approximately 500 pA using a range of magnifications from low to high.

Micro X-ray fluorescence (μXRF). Elemental maps (20 mm \times 10 mm) of dried SSRE cross sections were collected in air at 20 mbar using a Bruker M4 Tornado high performance μXRF spectrometer fitted with a Rh X-ray tube, poly-capillary optics and a Silicon Drift Detector. The poly-capillary optics allowed maps with a spot size diameter of 25 μm to be collected for Mo K_{α} . The mappings were performed with a pixel spacing of 25 μm with an acquisition time of 20 ms pixel⁻¹. The X-ray generator was operated at 50 kV and 599 μA using a 12.5 μM Al filter.

Electrochemical measurements

Electrochemical impedance spectroscopy. Measurements were performed in an aqueous 0.2 M KCl solution using each investigated reference electrode as the working electrode, coupled with a platinum wire counter electrode (CHI115, CH Instruments) and another saturated KCl Ag|AgCl reference electrode (1004 Series, Koslow Scientific). EIS measurements were performed in potentiostatic mode at open circuit potential with a 10 mV AC amplitude and a frequency range of 100 kHz to 10 mHz. Experiments were conducted in a Faraday cage using a PalmSens4 potentiostat. Replicate measurements were performed with three separate SSREs for each polymer composite type.

Potentiometric pH sensing. Measurements were performed in 2-electrode mode using a glass pH electrode (supplied by Ionode) coupled with each investigated reference electrode. The potential was monitored for 5 minutes in each buffer solution. The measurements were paused during buffer changes, and the electrodes were rinsed with deionised water and wiped dry with Kimwipe tissue before being immersed in the next buffer solution and continuing the measurements. A Faraday cage was needed for each test due to the high resistance of the glass pH electrode, and measurements were taken using a PalmSens4 potentiostat.

Cyclic voltammetry. Voltammograms were measured using an aqueous solution of 5 mM $[\text{Ru}(\text{NH}_3)_6]\text{Cl}_3$ with 0.2 M KCl

and scan rates between 25 mV s⁻¹ and 1 V s⁻¹. Scans were measured using a 1 mm glassy carbon working electrode (ET074, eDAQ) coupled with a platinum wire counter electrode (CHI115, CH Instruments) and the reference electrode under study. Measurements were taken using a PalmSens4 potentiostat.

Long-term potential measurements with sulphide. All tests measuring open circuit potential were maintained at *ca.* 25 °C using an incubator. Tests were conducted in 12.8 mM (1 g L⁻¹) Na₂S solution buffered at pH 9 to avoid sulphide losses as H₂S. The potential of each SSRE was measured simultaneously against the same SCE reference electrode (1001 Series, Koslow Scientific) by connecting the reference electrode cables from each potentiostat channel (VMP3, BioLogic). A double junction was used to protect the SCE and periodic maintenance was performed, however the SCE still drifted due to poisoning several times during the trials. The SSREs remained immersed in solution for the entire duration of the test.

Results and discussion

Impedance characterisation

The impedance of a reference electrode is an important parameter in determining the noise of an electrochemical experiment and the electrode's suitability for various electro-analytical techniques. To investigate the effect of suspended AgCl particles on electrode impedance, EIS was used to study both SSREs and compared to a typical liquid reference electrode, as shown in Fig. 2. These impedance values correspond to the same reference electrodes used for the potentiometric and voltammetric experiments presented later, however EIS was also repeated with replicate SSRE samples (see Fig. S2†). Each reference electrode demonstrated a mostly flat frequency response in the Bode plots, however higher impedance samples demonstrated larger phase angles in the high frequency range indicating a slowed voltage response.³¹ The mean impedance and standard deviation ($n = 3$) at 1.04 kHz were 16.3 ± 24.4 k Ω and 71.0 ± 51.7 k Ω for KCl and KCl/AgCl SSREs respectively. The large variability of impedance between electrodes of both SSREs may be due to differences in salt packing in the polymer matrix and Ag|AgCl wire positioning, however as it has been reported that most ionic conductivity in AgCl films occurs through micro-channels it may be expected that suspended AgCl particles in the polymer matrix will lower the conductivity of the material.³² All SSREs demonstrated impedances larger than those typically seen in liquid reference electrodes as the polymer matrix likely inhibits ionic conductivity, but these results are comparable to other resistance/impedance values reported for aqueous SSREs^{15,30} and lower than some reported ionic liquid based examples.^{3,33,34} The high impedance of these SSREs is likely to limit their use in fast CV and EIS because of noise and inductive effects respectively,^{31,35} but as we will show they provide comparable performance in potentiometry and CV experiments done at typical scan rates.



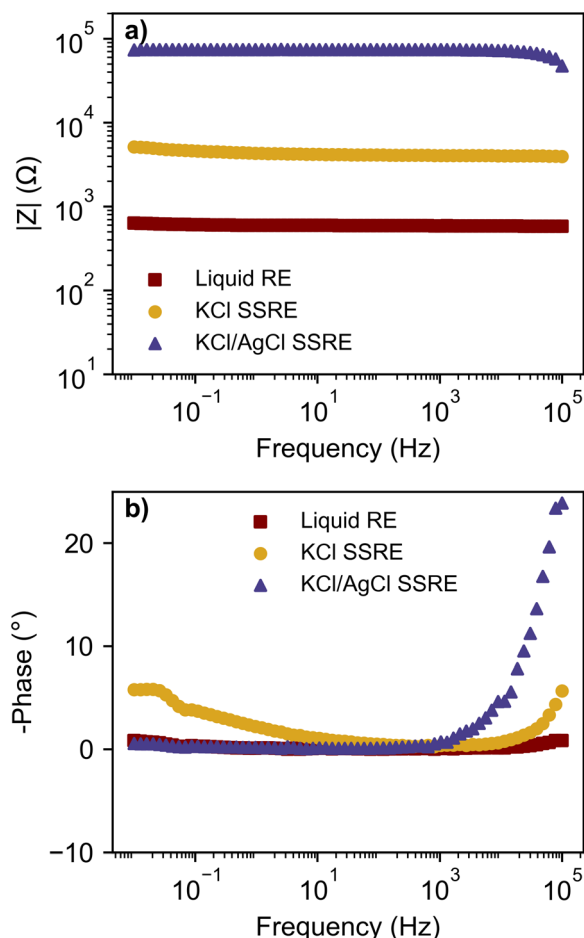


Fig. 2 Bode plots with (a) magnitude of impedance and (b) phase angle; comparing a liquid RE, KCl SSRE, and KCl/AgCl SSRE.

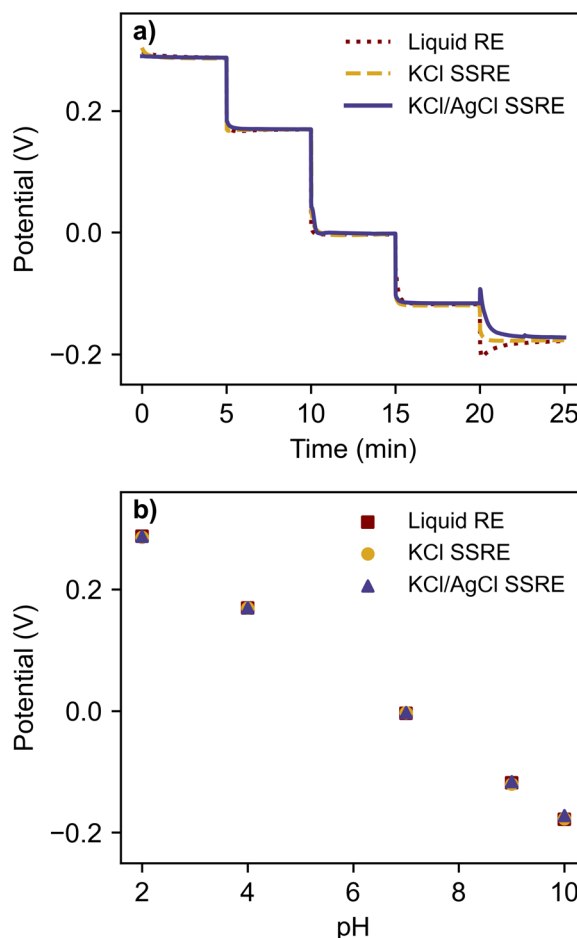


Fig. 3 (a) open circuit potentiometry with glass pH electrode in a series of pH buffers using a liquid RE, KCl SSRE, and KCl/AgCl SSRE, and (b) the resulting calibration curve.

Applications – potentiometric sensing and cyclic voltammetry

To compare the stability each reference electrode and demonstrate their use with a high impedance indicator electrode, each reference electrode was paired with a glass pH electrode and tested in a series of pH buffers, as summarised in Fig. 3 and Table 1.

Despite large differences in impedance, each reference electrode demonstrated equivalent performance, returning the expected linear calibration curves with similar Nernstian slopes and zero-point potentials.³⁶ The signal noise was less than 0.2 mV for all reference electrodes and buffer solutions. Transient spikes were seen in pH 10 buffer tests when using the liquid reference electrode and the KCl/AgCl SSRE, likely a result of rinsing and wiping the electrodes between buffer solutions rather than an effect of reference electrode impedance. These results demonstrate that the KCl/AgCl SSREs provided a stable reference potential across a wide pH range and would be suitable for use with a large range of indicator electrodes for potentiometric experiments, even those with high impedance.

The reference electrodes were also compared in CV experiments using the commonly used redox probe $[\text{Ru}(\text{NH}_3)_6]\text{Cl}_3$ at

Table 1 Summary of linear regression statistics from potentiometric testing using a glass pH electrode and three different reference electrodes

Reference electrode	Slope (mV pH^{-1})	Intercept (mV)	R^2
Liquid	-58.0	402.4	>0.9999
KCl SSRE	-57.8	401.2	>0.9999
KCl/AgCl SSRE	-57.4	400.7	>0.9999

scan rates between 25 mV s^{-1} and 1 V s^{-1} , as summarised in Fig. 4 and Table 2. The greater impedance of the KCl/AgCl SSRE did result in more signal noise compared to the other reference electrodes (Fig. S3–S5†), however this did not create significant differences in the analysis, even at the fastest scan rate of 1 V s^{-1} as seen in the voltammogram in Fig. 4a. Peak heights for the return peak were measured using a previous reported method of fitting a baseline to the diffusional current decay,³⁷ and showed good agreement between each reference electrode, demonstrating that near equivalent performance was achieved across the entire range of scan rates. The linear regression results for these peak currents vs. square root of



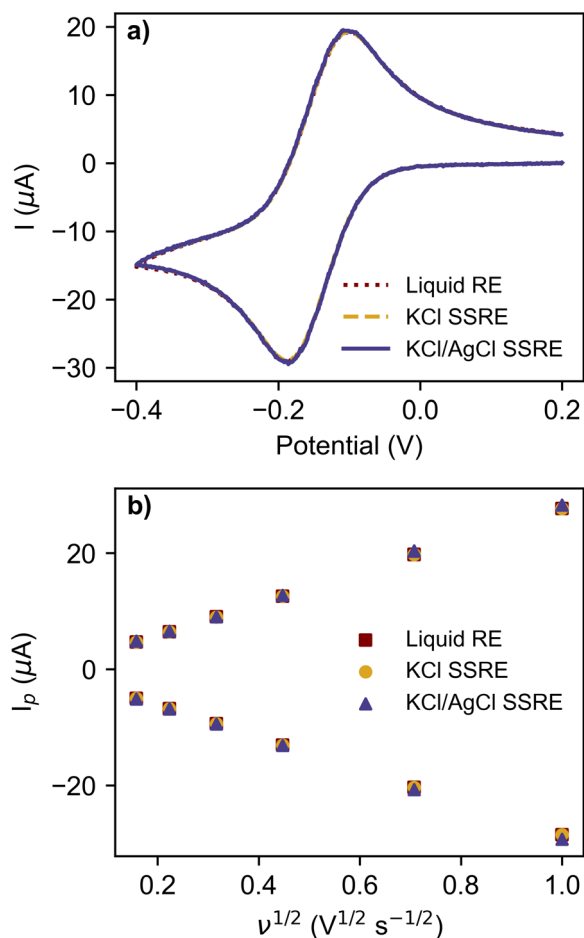


Fig. 4 (a) cyclic voltammograms using a 1 V s^{-1} scan rate with 5 mM $[\text{Ru}(\text{NH}_3)_6]\text{Cl}_3$ and 0.2 M KCl supporting electrolyte using a 1 mm diameter GC electrode with a liquid RE, KCl SSRE, and KCl/AgCl SSRE. (b) Peak currents (I_p) vs. square root of scan rate (ν) for voltammograms measured using various reference electrodes.

scan rate in Fig. 4b all demonstrated similar linearity, anodic, and cathodic slopes. Consequently, the diffusion coefficients calculated using these slopes with the Randles-Sevcik equation are all similar and close to previously reported values in the literature.^{38–41} These results demonstrate that the KCl/AgCl SSRE demonstrated comparable performance to a commercial liquid reference electrode, and should also prove suitable for chronoamperometric experiments. It should also be added that these datasets were analysed without any smoothing or post processing, however this may be useful in scenarios with a lower ratio of signal to noise.

Table 2 Summary of cyclic voltammetry analysis and linear regression

Reference electrode	Delta E_p range (mV)	Cathodic slope ($\mu\text{A s}^{1/2} \text{ V}^{-1/2}$)	Cathodic R^2	Anodic slope ($\mu\text{A s}^{1/2} \text{ V}^{-1/2}$)	Anodic R^2	$[\text{Ru}(\text{NH}_3)_6]^{3+}$ D ($\text{cm}^2 \text{ s}^{-1}$)	$[\text{Ru}(\text{NH}_3)_6]^{2+}$ D ($\text{cm}^2 \text{ s}^{-1}$)
Liquid	77–87	–28.757	0.9997	27.952	0.9998	7.41×10^{-6}	7.00×10^{-6}
KCl SSRE	76–84	–28.757	0.9997	27.935	0.9998	7.41×10^{-6}	6.99×10^{-6}
KCl/AgCl SSRE	77–83	–29.384	0.998	28.575	0.9998	7.74×10^{-6}	7.32×10^{-6}

Long-term sulphide resistance

To demonstrate the benefits of AgCl additive in the composite matrix, KCl SSREs and KCl/AgCl SSREs were immersed in a pH 9 buffer solution with 1 g L^{-1} Na_2S and left to condition *in situ* while their potentials were monitored vs. a SCE reference electrode, as summarised in Fig. 5 and Table 3. This solution was chosen to achieve sulphide concentrations well in excess of those typically found in wastewater systems and demonstrate the robustness of the KCl/AgCl composite SSRE.^{6,24} The pH buffer was used to prevent losses as H_2S over time, making HS^- the dominant sulphur species in these tests.⁴²

The SSREs were all monitored simultaneously against the same SCE reference electrode, and the literature value of -47 mV vs. SCE at 25°C was used to determine stabilisation and failure times.⁴³ Each SSRE was considered to have stabil-

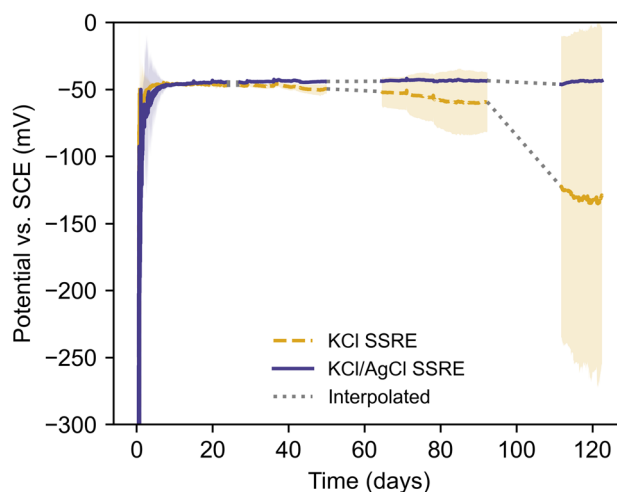


Fig. 5 long term potential monitoring of KCl SSRE and KCl/AgCl SSREs in 1 g L^{-1} Na_2S solution in pH 9 buffer. Mean potentials ($n = 5$) are reported with the coloured shading indicating the 95% confidence interval. Interpolated sections indicate periods where the conventional SCE reference suffered sulphide poisoning.

Table 3 Summary of OCP performance in 1 g L^{-1} Na_2S solution. Mean summary statistics ($n = 5$) reported with standard deviation

Reference electrode	Stabilisation time (days)	Failure time (days)	Working time (days)
KCl SSRE	2.0 ± 1.1	71.0 ± 25.1	69.1 ± 24.6
KCl/AgCl SSRE	3.1 ± 2.3	N/A	$>119.4 \pm 2.3$



used once their potential was within 5 mV of this value for 2 hours, and the failure time was recorded as the first time the potential deviated from this range after stabilisation. The working time was calculated as the difference between the last measurement before failure and the stabilisation time. Every SSRE adopted a potential within 2 mV of the literature value after stabilisation, however the KCl SSREs demonstrated varying stability with some exhibiting a negative potential drift before 20 days (Fig. S6†). These results demonstrate how sulphide species can significantly shorten the working life of a reference electrode as samples using this composite previously demonstrated a working life greater than 175 days in 1 M Na₂SO₄ solution without any electrode failures.³⁰ In contrast, no KCl/AgCl SSREs demonstrated this negative drift after stabilisation, and at *ca.* 120 days none had failed and all potentials were still close to the literature value (Fig. S7†). As such the variation in working time for KCl/AgCl SSREs in Table 3 is purely a result of differences in stabilisation time. Indeed, these electrodes were more stable in this environment than the commercial SCE used, which required consistent maintenance and occasional replacement due to poisoning over the course of the experiment. For this reason, affected periods of data have been omitted. It should be noted that the SSRE samples were continuously immersed during the entire experiment.

Fig. 6 summarises the results of μ XRF studies on cross-sections of SSREs after sulphide testing, and illustrates how suspended AgCl captures sulphide species and prevents reference electrode poisoning. The μ XRF distribution maps showed little difference in chlorine distribution between the two types of SSRE, suggesting a similar level of electrolyte loss. However, these experiments revealed significant differences in sulphur distri-

butions. For samples without AgCl suspended in the polymer, a dark precipitate formed around the tip of the embedded wire, whereas the KCl/AgCl samples had this precipitate concentrated at the open end of the polymer composite. This dark precipitate correlates with the sulphur distributions shown in μ XRF mapping, suggesting the formation of Ag₂S.^{25,26,28} Some signal was also seen correlating with the chlorine distribution, most notably when comparing Fig. 6c and e, suggesting the presence of trace sulphur impurities in the KCl. The low solubility product of Ag₂S also suggests that once this precipitate is formed the concentration of any aqueous sulphur species which could further diffuse through the polymer will be low.²⁹ These results demonstrate that the use of AgCl suspension could enable long-term continuous monitoring in high sulphide environments such as wastewater and groundwater. These cross-sections also demonstrate that although this design could be miniaturised, this will also result in a smaller electrolyte reservoir and would likely reduce the electrode's longevity.

Conclusions

A novel Ag|AgCl SSRE based on a KCl/AgCl polymer composite was developed which demonstrates exceptional long-term stability in challenging sulphide solutions. The impedance of the new SSRE was higher than a typical liquid filled reference electrode but comparable to other SSREs in the literature. This impedance was not limiting for potentiometric or voltammetric experiments and provided near-equivalent performance to the lower impedance liquid reference electrode. The KCl/AgCl SSREs demonstrated potentials close to the -47 mV *vs.* SCE expected from theory even after 120 days of continuous immersion in an aggressive sulphide solution, outlasting the KCl SSRE and commercial liquid RE without any maintenance. μ XRF measurements of SSRE cross-sections supported the hypothesis that suspended AgCl particles in the composite trap sulphide species, preventing their ingress and eventual poisoning of the Ag|AgCl electrode. This reference electrode should enable long-term continuous monitoring in wastewater and groundwater applications where sulphide is an ongoing challenge. More broadly, this work highlights a new approach for improving reference electrode longevity by capturing the interferant ions which cause poisoning, and suggests that similar composite materials could be explored for a range of other applications.

Author contributions

David S. Macedo: conceptualisation, methodology, formal analysis, investigation, writing – original draft, writing – review & editing, visualisation. Mikko Vepsäläinen: conceptualisation, methodology, writing – review & editing, supervision, project administration, visualisation. Theo Rodopoulos: writing – review & editing, supervision, visualisation. Stephen Peacock: formal analysis, visualisation. Conor F. Hogan: writing – review & editing, supervision, visualisation.

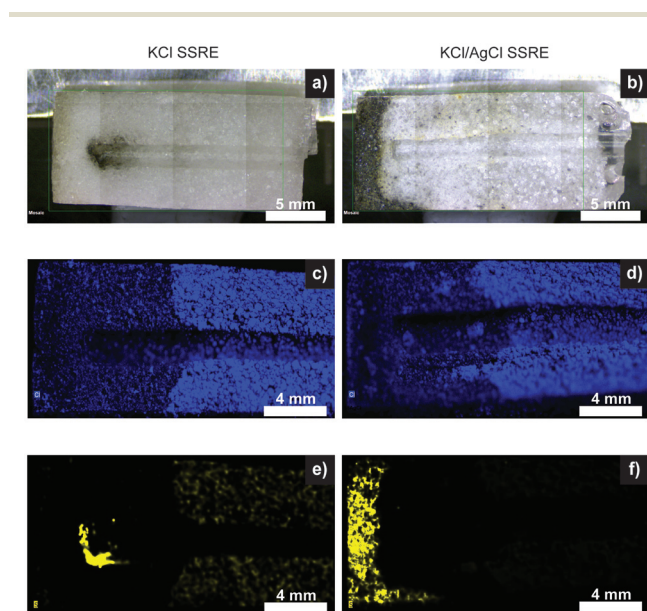


Fig. 6 KCl SSRE (left column) and KCl/AgCl SSRE (right column) cross section images (a and b) and μ XRF mapping of chlorine (c and d) and sulphur (e and f) after long term potential monitoring in 1 g L⁻¹ Na₂S solution.



Data availability

The data supporting this article have been included as part of the ESI.† The program and source code used for measuring return peak current in cyclic voltammetry experiments is available at <https://github.com/davedavedavem/diffusional-fitter>.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was funded by the Commonwealth Scientific and Industrial Research Organisation (CSIRO). The authors thank Luda Malishev for her assistance in electrode preparation, and Matthew Glenn for his assistance in SEM imaging.

References

- 1 P. Spitzer and S. Wunderli, in *Handbook of Reference Electrodes*, Springer Berlin Heidelberg, 2013, p. 77.
- 2 H. Kahlert, in *Electroanalytical Methods: Guide to Experiments and Applications*, ed. F. Scholz, Springer Berlin Heidelberg, Berlin, Heidelberg, 2002, pp. 261–278.
- 3 B. K. Troudt, C. R. Rousseau, X. I. N. Dong, E. L. Anderson and P. Bühlmann, *Anal. Sci.*, 2022, **38**, 71–83.
- 4 M. Cuartero, *Sens. Actuators, B*, 2021, **334**, 129635.
- 5 M. Cuartero and G. A. Crespo, *Curr. Opin. Electrochem.*, 2018, **10**, 98–106.
- 6 D. Despot, M. P. Fernández and M. Barjenbruch, *Water*, 2021, **13**, 1876.
- 7 A. Moretti, H. L. Ivan and J. Skvaril, *J. Water Process Eng.*, 2024, **60**, 105061.
- 8 P. Kruse, *J. Phys. D: Appl. Phys.*, 2018, **51**, 203002.
- 9 K. Nagy, K. Eine and K. Syverud, *J. Electrochem. Soc.*, 1997, **144**, 143–144.
- 10 H. J. Lee, U. S. Hong, D. K. Lee, J. H. Shin, H. Nam and G. S. Cha, *Anal. Chem.*, 1998, **70**, 3377–3383.
- 11 S. Gan, C. Liao, R. Liang, S. Du, L. Zhong, Y. Tang, T. Han, Y. Bao, Z. Sun, Y. Ma and L. Niu, *ACS Meas. Sci. Au*, 2022, **2**, 568–575.
- 12 Y. M. Wu and L. C. Chen, in *ISOEN 2019 – 18th International Symposium on Olfaction and Electronic Nose*, Proceedings, Institute of Electrical and Electronics Engineers Inc., 2019.
- 13 A. Lewenstam, B. Bartoszewicz, J. Migdalski and A. Kochan, *Electrochem. Commun.*, 2019, **109**, 106613.
- 14 M. Sophocleous and J. K. Atkinson, *Sens. Actuators, A*, 2017, **267**, 106–120.
- 15 R. C. Dawkins, D. Wen, J. N. Hart and M. Vepsäläinen, *Electrochim. Acta*, 2021, **393**, 139043.
- 16 M. Komoda, I. Shitanda, Y. Hoshi and M. Itagaki, *Electrochem. Commun.*, 2019, **103**, 133–137.
- 17 A. Moya, R. Pol, A. Martínez-Cuadrado, R. Villa, G. Gabriel and M. Baeza, *Anal. Chem.*, 2019, **91**, 15539–15546.
- 18 W. Gao, E. Zdrachek, X. Xie and E. Bakker, *Angew. Chem., Int. Ed.*, 2019, 1–6.
- 19 W. Gao, X. Xie and E. Bakker, *ACS Sens.*, 2020, **5**, 313–318.
- 20 E. L. Anderson, B. K. Troudt and P. Bühlmann, *ACS Sens.*, 2021, **6**, 2211–2217.
- 21 E. E. A. Robinson, B. K. Troudt and P. Bühlmann, *Anal. Chem.*, 2024, **96**, 2236–2243.
- 22 Z. Miao, M. L. Brusseau, K. C. Carroll, C. Carreón-Díazconti and B. Johnson, *Environ. Geochem. Health*, 2012, **34**, 539–550.
- 23 M. Ardelean, F. Manea, N. Vaszilcsin and R. Pode, *Anal. Methods*, 2014, **6**, 4775–4782.
- 24 M. Sherief and A. A. Hassan, *Water*, 2022, **14**, 791.
- 25 E. Lombi, E. Donner, S. Taheri, E. Tavakkoli, Å. K. Jämting, S. McClure, R. Naidu, B. W. Miller, K. G. Scheckel and K. Vasilev, *Environ. Pollut.*, 2013, **176**, 193–197.
- 26 R. Kaegi, A. Voegelin, B. Sinnet, S. Zuleeg, H. Siegrist and M. Burkhardt, *Sci. Total Environ.*, 2015, **535**, 20–27.
- 27 C. Levard, E. M. Hotze, G. V. Lowry and G. E. Brown, *Environ. Sci. Technol.*, 2012, **46**, 6900–6914.
- 28 Y. Yin, W. Xu, Z. Tan, Y. Li, W. Wang, X. Guo, S. Yu, J. Liu and G. Jiang, *Environ. Pollut.*, 2017, **220**, 955–962.
- 29 S. Licht, *J. Electrochem. Soc.*, 1988, **135**, 2971–2975.
- 30 D. S. Macedo, M. Vepsäläinen, D. Acharya, C. D. Wood, D. Wen, L. Thomson, S. Peacock, T. Rodopoulos and C. F. Hogan, *Electrochim. Acta*, 2021, **368**, 137636.
- 31 A. C. Lazanas and M. I. Prodromidis, *ACS Meas. Sci. Au*, 2023, **3**, 162–193.
- 32 H. Ha and J. Payer, *Electrochim. Acta*, 2011, **56**, 2781–2791.
- 33 X. V. Chen, A. Stein and P. Bühlmann, *ACS Sens.*, 2020, **5**, 1717–1725.
- 34 T. M. Galiullin, N. V. Pokhvisheva, A. V. Kalinichev and M. A. Peshkova, *Electroanalysis*, 2019, **31**, 1708–1718.
- 35 T. J. Smith and K. J. Stevenson, in *Handbook of Electrochemistry*, Elsevier, 2007, pp. 73–110.
- 36 A. Beard, M.-T. GmbH, S. Delina, J. Mettler-Toledo, S. Schmidt and H. Galster, *Ullmann's Encycl. Ind. Chem.*, 2023, 1–39.
- 37 D. S. Macedo, T. Rodopoulos, M. Vepsäläinen, S. Bajaj and C. F. Hogan, *Anal. Chem.*, 2024, **96**, 1530–1537.
- 38 Y. Wang, J. G. Limon-Petersen and R. G. Compton, *J. Electroanal. Chem.*, 2011, **652**, 13–17.
- 39 Á. Molina, E. Laborda, E. I. Rogers, F. Martínez-Ortiz, C. Serna, J. G. Limon-Petersen, N. V. Rees and R. G. Compton, *J. Electroanal. Chem.*, 2009, **634**, 73–81.
- 40 T. L. Ferreira, T. R. L. C. Paixão, E. M. Richter, O. A. El Seoud and M. Bertotti, *J. Phys. Chem. B*, 2007, **111**, 12478–12484.
- 41 K. J. Levey, M. A. Edwards, H. S. White and J. V. Macpherson, *Phys. Chem. Chem. Phys.*, 2023, **25**, 7832–7846.
- 42 A. E. Lewis, *Hydrometallurgy*, 2010, **104**, 222–234.
- 43 A. J. Bard, L. R. Faulkner and H. S. White, *Electrochemical Methods: Fundamentals and Applications*, John Wiley & Sons, Ltd, 3rd edn, 2022.

