Due to rapid industrialisation the atmosphere is polluted by acidic gases like $\text{SO}_2$, $\text{H}_2\text{~S}$, $\text{NO}_2$ etc, which increase the corrosivity of atmosphere. For continuous monitoring of the atmospheric corrosivity a probe and coulometer have been designed. From the measured coulombs the corrosivity of atmosphere can be monitored. This paper presents the results of the performance of the monitor.

Keywords: Atmospheric corrosivity, probe, coulometer.

INTRODUCTION

Material degradation due to corrosion is a severe problem. The economics of this problem forces one to look for a better monitoring and good remedial measure. The atmospheric corrosion is an electrochemical phenomenon with an electrochemical reaction which is highly complicated [1-3]. Unlike the general corrosion which occurs in an aqueous phase, the atmospheric corrosion takes place when there exists an electrolyte layer over the surface exposed to the atmosphere. This electrolyte layer normally originates from the condensed phase of moisture which is extremely thin (approximately 1 μm). Factors like temperature, humidity, and the other pollutants present in the air contribute significantly to the corrosion due to this thin electrolyte layer. Due to rapid industrialisation, the atmosphere gets polluted by the flue gases containing acidic and toxic gases like $\text{SO}_2$, $\text{H}_2\text{~S}$, $\text{NO}_2$ etc. These pollutants which are soluble in the moisture of the atmosphere enhances the rate of corrosion.

The loss of metal through the atmospheric corrosion is a direct function of time $t_i$ during which the thin aqueous film persists over the surface as given by

$$W = r \times t_i$$

where ‘r’ is the average corrosion rate.

Since the conventional means of monitoring corrosion requires the electrodes to be immersed in electrolyte which do not mimic the situation of atmospheric corrosion, extensive attempts [4-8] have been reported in designing newer experimental setups to assess the atmospheric corrosion and to forecast its corrosion rate. Among all the new experiments developed, the most popular one is the use of monitoring the galvanic current flowing through a galvanic couple.

In this paper a new electrode arrangement and a five digit digital coulometer have been reported. The use of this setup for the atmospheric corrosion has been tested with the acidic vapour condensation setup.

Circuit description

The principle of working of the environmental corrosivity monitor can be explained with the help of a schematic block diagram shown in Fig. 1. The block diagram consists of a

![Fig. 1: Block diagram of environmental corrosivity monitor](image-url)
The arrangement of the electrochemical probe

Current-to-voltage converter (CVC)

The schematic circuit diagram of CVC is shown in Fig. 3. The output voltage $V_o$ is proportional to the input current $I_{in}$ and is given by:

$$V_o = R_t \times I_{in}$$

This circuit is used for input current ranges up to 1 A, and the output is fed to the input of the unipolar generator.

Unipolar generator (UPG)

IC 2 and IC 3 constitutes the unipolar generator. When input is positive, it results in a negative output at IC 2 due to rectification. This negative signal is inverted by IC 3 and a positive signal is obtained at the output. When the input is negative, it is directly inverted by IC 3. Thus the output of UPG is always positive. The voltage to frequency converter responds only for positive input voltages and therefore this stage is needed.

Voltage-to-frequency converter (VFC)

In this application, the XR-4151 (IC 4) functions as a stand alone VFC operating on a single positive power supply [9]. The functional block diagram of XR-4151 is shown in Fig. 4. It consists of a voltage comparator, a one shot, and a precision switched current source. The output of the UPG is applied at pin 7 of the VFC and the VFC gives a periodic square wave. The voltage comparator compares the input voltage to the voltage at pin 6. If the input voltage is higher, the comparator will fire the one-shot monostable multivibrator. The output of the one-shot is connected to both the logic output and the precision switched current source. During the one-shot period $T$, the logic output will go low and the current source will turn on with current $I$. At the end of the one-shot period the logic output will go high and the current source has injected an amount of charge $Q = I_o \times T$ into the network $R_o - C_B$. If this injecting charge does not increase the voltage $V_B$ such that $V_B > V_T$, the comparator again fires the one-shot and the current source injects another lump of charge $Q$, into the $R_B - C_B$ network. This process continues until $V_B$ is again equal to $V_T$. This completes one cycle. The VFC will now run in a steady state mode. The current source dumps lumps of charge into the capacitor $C_B$ at the rate fast enough to keep $V_B > V_T$. Since the discharge rate of capacitor $C_B$ is proportional to $V_B/R_B$, the frequency at which the system runs will be proportional to the input voltage on pin 7 and is given by:

$$F_o \propto V_1$$

where $K = 0.486 R_e/(R_B R_o C_o)$.

The input voltage range is from 0 to $+10$ V, and the output frequency is from 0 to 10 kHz.

Counter

The frequency output from VFC is fed to the count input of the counter ICM 7217M (IC 7) through an AND gate, and a clock signal from the timer NE 555 (IC 5) is also applied to the gate that desires the counting time. The counter 7217A is a four digit LED display programmable up/down counter.
It directly drives a multiplexed seven segment LED display. This circuit provides three main outputs; a CARRY/BORROW output which allows for direct cascading of counters, a zero output which indicates when the count is zero and an equal output which indicates when the count is equal to the value contained in the registers. The up/down counting is decided by the control signal from the polarity detector (IC 6). A manual switch is also provided to reset the count so that the system can start counting from the beginning. By cascading one more counter chip (IC 8) counting is extended to five digits. The four output lines of the decade counter (IC 8) are directly connected to the decoder (IC 9) to drive the fifth display.

By properly calibrating and scaling the input current to get the corresponding digital output to the counter to read out in coulombs is achieved.
EXPERIMENTAL

A co-axial galvanic cell consisting of inner copper and outer steel electrode of area $2 \text{ cm}^2$ each separated by a very thin Teflon layer mounted in a Teflon body with electrical contacts taken in the other side was used. The unexposed side of the probe was moulded with epoxy resin to protect the electrical contacts.

This probe was kept hanging in a closed glass vessel containing 50% and 100% HCl.

RESULTS AND DISCUSSION

The results of the investigation is presented in Table I which shows the charge flow for the 50% and 100% HCl vapour. It is evident that the rate of increase and the coulombs increase for 100% is comparable to that of 50%. This confirms the severity of the 100% HCl which is very well known. The measurements after three hours of keeping the probe inside the 50% instantaneous results, is still higher than 100% concentration clearly points out that exposure time, i.e. the actual time of existence of aqueous layer ($t$) has a critical role over the rate of corrosion.

Specifications

Maximum input current - 1 A, maximum counts - 99999, output - 99.999 C and energy source - 230 V, 50 Hz.

CONCLUSION

The probe and the circuitry designed during this investigation can be successfully used to monitor the atmospheric corrosion. A light modification in the probe assembly with an air sucking facility will help in using this instrument as an environmental corrosivity monitor in the industries.

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