



NRL Report 8107

Mathematical Analysis of Circular Corrosion Cells Having Unequal Polarization Parameters

E. MCCAFFERTY

Metals Performance Branch Engineering Materials Division

August 5, 1977



NAVAL RESEARCH LABORATORY Washington, D.C.

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unequal anodic and cathodic polarization parameters is not related in a simple manner to the distribution curves for equal parameters. For bulk electrolyte the value of the electrode potential across the surface depends on whether the system is under anodic, cathodic, or mixed control. The current distribution is the more uniform for combinations of more polarizable electrodes. In thin-layer electrolytes there is a geometry effect in which electrode polarization and current flow are concentrated near the anode/ cathode juncture.

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MATHEMATICAL ANALYSIS OF CIRCULAR CORROSION CELLS HAVING UNEQUAL POLARIZATION PARAMETERS

INTRODUCTION

In many corrosion reactions the anode and cathode are spatially localized. This may occur on different surfaces, as in the galvanic corrosion of dissimilar metals, or on different parts of the same surface, as with localized geometries such as crevices. Moreover the corroding system often has a coplanar concentric circular geometry. As pointed out earlier [1, 2], examples include some instances of pitting [3], crevice corrosion under Orings or washers [4], and corrosion under barnacles [5], under tubercules of corrosion products [6], or under dust particles in condensed moisture films [7].

In all such cases there is a potential difference between the central anode and the disk-shaped cathode surrounding it. This potential difference may arise from heterogeneities in the solid phase (such as dissimilar metals, inclusions in a base metal, or discontinuities in protective films) or from heterogeneities in the liquid phase (such as differential aeration as in crevices). Thus there is a distribution of both electrode potential and local current density as one moves radially from the center of the anode out toward the far edge of the cathode.

Gal-Or, Raz, and Yahalom [8] have mathematically treated systems of coplanar concentric circular corrosion cells. These authors analyzed the effect of various system parameters on the total current, and more recently McCafferty [1, 2] has evaluated the distribution of potential and current across such cells. These treatments essentially extended to cylindrical geometries the model developed by Waber and coworkers [9-12] in a series of publications treating semi-infinite parallel electrodes.

Two central features in the Waber model are that the anode and cathode obey linear polarization kinetics over an extended potential range and that the anodic and cathodic slopes are equal. Whereas the first assumption often holds in experiments, the second assumption rarely holds, because the anode is generally far less polarizable than the cathode.

The case of unequal anodic and cathodic lineæ by Kennard and Waber [13] for semi-infinite strips electrolyte.

tion has been solved recently el electrodes under bulk

This report extends the Waber model of linear corrosion kinetics to circular systems with unequal polarization parameters. Equations are derived for potential and current distributions and for the total anodic current, and generalized calculations are made. Comparisons with experimental results will be made elsewhere.

Manuscript submitted February 7, 1977.

DESCRIPTION OF THE MODEL

The Corrosion Cell

The cell geometry is shown in Fig. 1. The anode and cathode outer edges are coplanar concentric circles of radii a and c respectively. The electrolyte thickness b is allowed to approach infinity for bulk electrolyte.

Linear Polarization

The cell potentials are shown in Fig. 2a, and stylized polarization curves are shown in Figs. 2b and 2c. Following Wagner [14] and Waber [9-13], an important feature of the model is that the polarization curves are linear in E vs *i* over an extended range. As pointed out by Kennard and Waber [13], if the plots are linear over only a portion of the curve, tangent approximations can be drawn. Thus the open-circuit potentials E_a^o and E_c^o are replaced by the intersections $E_a^{o'}$ and $E_c^{o'}$ respectively of the tangent lines with the potential axis, as shown in Fig. 2c.



Fig. 1-The corrosion cell

2





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Fig. 2c—Linear approximations to the polarization curves. The extrapolated values $E_a^{o'}$ and $E_c^{o'}$ replace E_a^{o} and E_c^{o} respectively.

The linearized polarization curves are characterized by the Wagner polarization parameters

$$\mathcal{L}_a = \sigma \left| \frac{dE}{di} \right|_a \tag{1}$$

for the anode and

$$\mathcal{L}_{c} = \left. \sigma \right| \frac{dE}{di} \bigg|_{c} \tag{2}$$

for the cathode, where σ is the electrolyte conductivity. The parameters \mathcal{L}_a and \mathcal{L}_c have the dimensions of length (cm), and $\mathcal{L}_a \neq \mathcal{L}_c$ in the present treatment.

The assumption of linear polarization over an extended potential range has been observed to be a reasonable approximation in a number of instances. For example, steels in aerated neutral to basic solutions, with or without chloride, displayed both anodic and cathodic plots which were approximately linear over an extended range [15]. Additional examples include the behavior of copper/steel couples in distilled water [16], the corrosion of tin in citrate solutions [17], and the corrosion of bare and coated aluminum in chloride solutions [18]. Other examples involve specialized geometries, such as the pitting of aluminum [19], or specialized conditions, such as the dissolution of mild steel at high anodic overpotentials in concentrated electrolytes [20]. On the cathodic side the reduction of oxygen on nickel in dilute H_2SO_4 [21] and of silver in KOH [22] display linear regions. Polarization curves for a variety of metals in thin-layer electrolytes [23] display linear regions over at least part of the potential ranges for both anodic and cathodic processes.

In some cases the linearity may be attributed predominantly to resistance polarization, caused either by iR drops through the solution or by ohmic films on the electrode surface. As pointed out by Stern and Geary [24], however, sometimes the combined effects of concentration polarization plus ohmic drops interfere with activation polarization processes so that a very short Tafel region is observed. Such cases often give straight-line segments in E vs *i*.

At this point it should be clear that the model invokes linearity over an extended potential range and not merely in the pre-Tafel region near the corrosion potential, where the usual Stern and Geary [24] linear relation is valid.

MATHEMATICAL ANALYSIS FOR BULK ELECTROLYTE

The electrostatic potential P(x, y, z) is given by Laplace's equation

$$\nabla^2 P(x, y, z) = 0, \tag{3}$$

provided that there are no concentration gradients in the solution, the solution is electroneutral, and there are no sources or sinks of ions in the electrolyte [25].

With the circular geometry it is convenient to rewrite Eq. (3) in cyclindrical coordinates using the usual transformations $x = r \cos \theta$ and $y = r \sin \theta$. The result is

$$\frac{\partial^2 P(r,z)}{\partial r^2} + \frac{1}{r} \frac{\partial P(r,z)}{\partial r} + \frac{\partial^2 P(r,z)}{\partial z^2} = 0, \qquad (4)$$

where the potential P is independent of the angle θ . The general approach is to solve for P(r, z) in Eq. (4) subject to appropriate boundary conditions and then to evaluate the local current density i(r, 0) from Ohm's law for electrolytes:

$$i(r, 0) = -\sigma \left[\frac{\partial P(r, z)}{\partial z} \right]_{z=0},$$
(5)

where σ is the electrolyte conductivity.

Boundary Conditions

The boundary conditions have been discussed in some detail in a previous report [1]. In brief there is no current flow across the symmetry line r = 0, nor across the cathode outer boundary r = c. Thus

$$\left[\frac{\partial P(r,z)}{\partial r}\right]_{r=0} = 0 \tag{6}$$

and

$$\left[\frac{\partial P(r,z)}{\partial r}\right]_{r=c} = 0.$$
⁽⁷⁾

Also, the potential must be bounded at the upper physical boundary of the electrolyte, so that

$$\lim_{z \to \infty} P(r, z) < M, \tag{8}$$

where M is some finite number.

The general solution to Eq. (4) subject to the boundary conditions of Eqs. (6) through (8) is [1, 2, 8]

$$P(r, z) = C_0 + \sum_{n=1}^{\infty} C_n J_0(\lambda_n r) e^{-\lambda_n z}, \qquad (9)$$

where C_0 and C_n are coefficients to be evaluated later, J_0 is the Bessel function of order zero, and $\lambda_n = x_n/c$, in which the x_n are the zeros of $J_1(x) = 0$.

Linear Polarization

The remaining boundary condition relates the electrode potential E(r, 0) along the metal surface vs some standard reference electrode to the electrostatic potential P(r, 0) within the electrolyte but "just outside" [26] the electrode surface. If E_a and E_c are the potentials of the polarized anode and cathode respectively, at any current density, then

$$V' - P(r, 0) = E(r, 0), \qquad (10)$$

where V' is a constant which includes the various differences in electrostatic potential across the extra interfaces introduced in the measurement of a potential difference across the metal/solution interface of interest [27]. Equation (10) is developed in Appendix A.

For the anodic branch in Fig. 2b,

slope =
$$\frac{i(r, 0) - 0}{E(r, 0) - E_a^o} = \left| \frac{di}{dE} \right|_a$$
, (11)

which after rearranging becomes

$$E(r, 0) = E_a^o + i(r, 0) \left| \frac{dE}{di} \right|_a.$$
 (12)

Substitution of Eq. (5) in Eq. (12) gives

$$E(r, 0) = E_a^o - \sigma \left| \frac{dE}{di} \right|_a \left[\frac{\partial P(r, z)}{\partial z} \right]_{z=0}$$
(13)

Use of the Wagner polarization parameter as defined in Eq. (1) gives:

$$E(r, 0) = E_a^o - \mathcal{L}_a \left[\frac{\partial P(r, z)}{\partial z} \right]_{z=0}.$$
 (14)

Substitution of Eq. (14) in Eq. (10) gives

$$P(r, 0) - \mathcal{L}_a \left[\frac{\partial P(r, z)}{\partial z} \right]_{z=0} = V' - E_a^o, \quad 0 \leq r < a.$$
(15a)

A similar expression holds for the cathode:

$$P(r, 0) - \pounds_c \left[\frac{\partial P(r, z)}{\partial z} \right]_{z=0} = V' - E_c^0, \quad a < r \le c.$$
(15b)

Equations (15a) and (15b) are the final boundary conditions required. The indeterminate constant V' will vanish in the final forms of the expressions to be derived but will be carried along for mathematical completeness.

Evaluation of the Coefficients Cn

The boundary conditions in Eqs. (15a) and (15b) are used to determine the coefficients C_n appearing in Eq. (9). The approach is to evaluate Eqs. (15a) and (15b) using the general expression for P(r, z) and then to solve the two simultaneous equations. The reader wishing to avoid the mathematical details can skip to Eq. (35).

The general expression for P(r, z) was given earlier by Eq. (9). Use of Eq. (9) in (15a) gives

$$C_0 + \sum_{n=1}^{\infty} C_n (1 + f_a \lambda_n) J_0(\lambda_n r) = V' - E_a^o, \quad 0 \le r < a.$$
 (16)

If this equation is multiplied through by $rJ_0(\lambda_m r)$ and integrated over the domain of applicability (from r = 0 to r = a), then

$$C_{0} \int_{r=0}^{a} r J_{0}(\lambda_{m} r) dr + \int_{r=0}^{a} \sum_{n=1}^{\infty} C_{n}(1 + \mathcal{L}_{a}\lambda_{n}) r J_{0}(\lambda_{n} r) J_{0}(\lambda_{m} r) dr$$
$$= (V' - E_{a}^{o}) \int_{r=0}^{a} r J_{0}(\lambda_{m} r) dr.$$
(17)

The first and third integrals can be evaluated from a standard recursion formula for Bessel functions [28]; that is

$$\frac{d}{dx}\left[xJ_1(x)\right] = xJ_0(x), \tag{18}$$

which, upon appropriate variable change and integration, gives

$$\int r J_0(\lambda r) dr = \frac{1}{\lambda} r J_1(\lambda r).$$
(19)

The second integral to be called I_2 , is

$$I_2 \equiv \int_{r=0}^{a} \sum_{n=1}^{\infty} C_n (1 + \mathcal{L}_a \lambda_n) r J_0(\lambda_n r) J_0(\lambda_m r) dr \qquad (20)$$

or

$$I_2 = \sum_{n=1}^{\infty} C_n (1 + \mathcal{L}_a \lambda_n) \int_{r=0}^a r J_0(\lambda_n r) J_0(\lambda_m r) dr.$$
(21)

The summation can be split into two cases: n = m, and $n \neq m$. Thus

$$I_{2} = C_{m}(1 + \pounds_{a}\lambda_{m}) \int_{r=0}^{a} rJ_{0}^{2}(\lambda_{m}r) dr$$
$$+ \sum_{\substack{n=1\\n \neq m}}^{\infty} C_{n}(1 + \pounds_{a}\lambda_{n}) \int_{r=0}^{a} rJ_{0}(\lambda_{n}r)J_{0}(\lambda_{m}r) dr.$$
(22)

Substitution of Eqs. (19) and (22) back into Eq. (17) thus yields

$$C_{0} \frac{1}{\lambda_{m}} aJ_{1}(\lambda_{m}a) + C_{m}(1 + \pounds_{a}\lambda_{m}) \int_{r=0}^{a} rJ_{0}^{2}(\lambda_{m}r) dr$$

$$+ \sum_{\substack{n=1\\n \neq m}}^{\infty} C_{n}(1 + \pounds_{a}\lambda_{n}) \int_{r=0}^{a} rJ_{0}(\lambda_{n}r)J_{0}(\lambda_{m}r) dr$$

$$= (V' - E_{a}^{0}) \frac{1}{\lambda_{m}} aJ_{1}(\lambda_{m}a). \qquad (23)$$

The second integral in Eq. (23) is a Lommel integral [29]:

$$(\alpha^2 - \beta^2) \int_{x_1}^{x_2} x J_n(\alpha x) J_n(\beta x) dx = \left[x \left\{ \beta J_n(\alpha x) J_n'(\beta x) - \alpha J_n'(\alpha x) J_n(\beta x) \right\} \right]_{x_1}^{x_2}$$
(24)

where the primes denote differentiation with respect to the whole argument and not just x. Thus, when $n \neq m$,

$$\int_{0}^{a} r J_{0}(\lambda_{n}r) J_{0}(\lambda_{m}r) dr = \frac{a}{\lambda_{n}^{2} - \lambda_{m}^{2}} \left[\lambda_{n} J_{1}(\lambda_{n}a) J_{0}(\lambda_{m}a) - \lambda_{m} J_{0}(\lambda_{n}a) J_{1}(\lambda_{m}a) \right].$$
(25)

When n = m, the integral becomes $\int r J_0^2(\lambda_m r) dr$, which is the remaining integral to be evaluated in Eq. (23). However, when n = m, the right-hand side of Eq. (25) gives 0/0, so that l'Hospital's rule must be used. In this case the numerator and denominator are differentiated with respect to λ_n , and then λ_n is allowed to approach λ_m . The result (omitting several steps) is

$$\int_{r=0}^{a} r J_{0}^{2}(\lambda_{m}r) dr = \frac{a^{2}}{2} \left[J_{0}^{2}(\lambda_{m}a) + J_{1}^{2}(\lambda_{m}a) \right].$$
(26)

Use of Eqs. (25) and (26) in Eq. (23) gives

$$C_{0} \frac{1}{\lambda_{m}} a J_{1}(\lambda_{m} a) + C_{m}(1 + \pounds_{a} \lambda_{m}) \frac{a^{2}}{2} \left[J_{0}^{2}(\lambda_{m} a) + J_{1}^{2}(\lambda_{m} a) \right]$$

$$+ \sum_{\substack{n=1\\n \neq m}}^{\infty} C_{n}(1 + \pounds_{a} \lambda_{n}) \frac{a}{\lambda_{n}^{2} - \lambda_{m}^{2}} \left[\lambda_{n} J_{1}(\lambda_{n} a) J_{0}(\lambda_{m} a) - \lambda_{m} J_{0}(\lambda_{n} a) J_{1}(\lambda_{m} a) \right]$$

$$= (V' - E_{a}^{o}) \frac{a}{\lambda_{m}} J_{1}(\lambda_{m} a). \qquad (27)$$

Equation (27) is one of the two simultaneous equations to be solved for the set C_n . The second equation follows from the boundary condition on the cathode given in Eq. (15b). The approach is the same as has just been completed. P(r, 0) and $\partial P(r, z)/\partial z$ at z = 0 are evaluated from Eq. (9), so that Eq. (15b) becomes

$$C_0 + \sum_{n=1}^{\infty} C_n (1 + f_c \lambda_n) J_0(\lambda_n r) = V' - E_c^o, \quad a < r \le c.$$
 (28)

Again the equation is multiplied by $rJ_0(\lambda_m r) dr$ and integrated over the appropriate limits, which in this case are from r = a to r = c:

$$C_0 \int_{r=a}^{c} r J_0(\lambda_m r) dr + \int_{r=a}^{c} \sum_{n=1}^{\infty} C_n (1 + \mathcal{L}_c \lambda_n) r J_0(\lambda_n r) J_0(\lambda_m r) dr$$
$$= (V' - E_c^o) \int_{r=a}^{c} r J_0(\lambda_m r) dr.$$
(29)

The first and third integrals can be evaluated using Eq. (19):

$$\frac{C_0}{\lambda_m} \left[cJ_1(\lambda_m c) - aJ_1(\lambda_m a) \right] + \sum_{n=1}^{\infty} C_n (1 + \mathcal{L}_c \lambda_n) \int_{r=a}^{c} rJ_0(\lambda_n r) J_0(\lambda_m r) dr$$
$$= \frac{V' - E_c^0}{\lambda_m} \left[cJ_1(\lambda_m c) - aJ_1(\lambda_m a) \right]. \tag{30}$$

9

By definition $\lambda_m c = x_m$ and $J_1(x_m) = 0$. Again the summation can be split into two cases, so that Eq. (30) becomes

$$-\frac{C_0}{\lambda_m} a J_1(\lambda_m a) + C_m (1 + \mathcal{L}_c \lambda_m) \int_{r=a}^c r J_0^2(\lambda_m r) dr$$

$$+ \sum_{\substack{n=1\\n \neq m}}^{\infty} C_n (1 + \mathcal{L}_c \lambda_n) \int_{r=a}^c r J_0(\lambda_n r) J_0(\lambda_m r) dr$$

$$= -\frac{(V' - E_c^0)}{\lambda_m} a J_1(\lambda_m a).$$
(31)

The integrals in Eq. (31) can be evaluated as before from Eq. (24). A simpler approach is to add and subtract $\int_0^a r J_0^2(\lambda_m r) dr$ to the second term and to add and subtract $\int_0^a r J_0(\lambda_n r) J_0(\lambda_m r) dr$ within the summation sign: Then Eq. (31) becomes

$$-\frac{C_0}{\lambda_m} a J_1(\lambda_m a) + C_m (1 + \mathcal{L}_c \lambda_m) \left[\int_{r=0}^c r J_0^2(\lambda_m r) dr - \int_{r=0}^a r J_0^2(\lambda_m r) dr \right]$$

+
$$\sum_{\substack{n=1\\n \neq m}}^{\infty} C_n (1 + \mathcal{L}_c \lambda_n) \left[\int_{r=0}^c r J_0(\lambda_n r) J_0(\lambda_m r) dr - \int_{r=0}^a r J_0(\lambda_n r) J_0(\lambda_m r) dr \right]$$

=
$$-\frac{(V' - E_c^0)}{\lambda_m} a J_1(\lambda_m a).$$
(32)

The integrals involving the entire interval from r = 0 to r = c are the usual orthogonality relations [30],

$$\int_{r=0}^{c} r J_{0}(\lambda_{n} r) J_{0}(\lambda_{m} r) dr = \begin{cases} 0, n \neq m, \\ \frac{c^{2}}{2} [J_{0}^{2}(\lambda_{m} c)], n = m, \end{cases}$$
(33)

and the integrals from r = 0 to r = a have already been evaluated per Eqs. (25) and (26), so that Eq. (32) reduces to

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$$-\frac{C_{0}}{\lambda_{m}}aJ_{1}(\lambda_{m}a) + C_{m}(1 + \mathcal{L}_{c}\lambda_{m})\left\{\frac{c^{2}}{2}J_{0}^{2}(\lambda_{m}c) - \frac{a^{2}}{2}[J_{0}^{2}(\lambda_{m}a) + J_{1}^{2}(\lambda_{m}a)]\right\}$$

$$+\sum_{\substack{n=1\\n\neq m}}^{\infty}C_{n}(1 + \mathcal{L}_{c}\lambda_{n})\left\{-\frac{a}{\lambda_{n}^{2} - \lambda_{m}^{2}}[\lambda_{n}J_{1}(\lambda_{n}a)J_{0}(\lambda_{m}a) - \lambda_{m}J_{0}(\lambda_{n}a)J_{1}(\lambda_{m}a)]\right\}$$

$$=-\frac{(V' - E_{c}^{0})}{\lambda_{m}}aJ_{1}(\lambda_{m}a).$$
(34)

Equations (34) and (27) are thus two simultaneous equations in C_0 and C_n . Addition of the two equations eliminates C_0 . The result, after considerable algebra, is

$$C_{m}\left[\left(1+f_{c}\frac{x_{m}}{c}\right)\frac{J_{0}^{2}(x_{m})}{2}+(f_{a}-f_{c})\left(\frac{x_{m}}{c}\right)\left(\frac{a}{c}\right)^{2}A_{m}\right]+\frac{a}{c^{2}}(f_{a}-f_{c})\sum_{\substack{n=1\\n\neq m}}^{\infty}C_{n}W_{nm}$$

$$=-\frac{(E_{a}^{o}-E_{c}^{o})}{2}\left(\frac{a}{c}\right)J_{1}\left(x_{m}\frac{a}{c}\right),$$
(35)

$$x_m \quad (c)^{\sigma_1} (a^m c),$$

where A_m and W_{nm} are defined by

$$A_m = \frac{1}{2} \left[J_0^2 \left(x_m \frac{a}{c} \right) + J_1^2 \left(x_m \frac{a}{c} \right) \right]$$
(35a)

and

$$W_{nm} = \frac{x_n^2}{x_n^2 - x_m^2} \left[J_1\left(x_n \frac{a}{c}\right) J_0\left(x_m \frac{a}{c}\right) - \frac{x_m}{x_n} J_0\left(x_n \frac{a}{c}\right) J_1\left(x_m \frac{a}{c}\right) \right].$$
(35b)

Equation (35) thus generates a series of equations, say k of them, where m is fixed in turn from 1 through k. These k equations are solved simultaneously to give the coefficients C_n from n = 1 to k.

The indeterminate constant V' cancels out in the generation of the set of C_n . Also, when $\mathcal{L}_a = \mathcal{L}_c = \mathcal{L}$, Eqs. (35a) and (35b) reduce to the previous case [1, 2]:

$$C_{n} = -\frac{(E_{a}^{o} - E_{c}^{o})\left(\frac{a}{c}\right)J_{1}\left(x_{n}\frac{a}{c}\right)}{x_{n}\left(1 + \pounds\frac{x_{n}}{c}\right)\frac{J_{0}^{2}(x_{n})}{2}},$$
(36)

where the dummy variable m has been replaced by the more general n.

Evaluation of the Coefficient Co

The remaining coefficient C_0 can be most conveniently evaluated by going back to the original pair of equations: Eqs. (16) and (28). The approach is to multiply through in both equations by rdr and then to integrate over the appropriate limits. The operations are straightforward, so that it is not necessary to detail the proof here. The resulting two equations are

$$\frac{a^2}{2} C_0 + \sum_{n=1}^{\infty} C_n (1 + \mathcal{L}_a \lambda_n) \frac{a}{\lambda_n} J_1(\lambda_n a) = (V' - E_a^o) \frac{a^2}{2}$$
(37)

and

$$\frac{(c^2 - a^2)}{2}C_0 - \sum_{n=1}^{\infty} C_n (1 + \pounds_c \lambda_n) \frac{a}{\lambda_n} J_1(\lambda_n a) = (V' - E_c^o) \left(\frac{c^2 - a^2}{2}\right)$$
(38)

Adding Eqs. (37) and (38) and solving for C_0 gives

$$C_0 = V' - \left(\frac{a}{c}\right)^2 E_a^o - \left(\frac{c^2 - a^2}{c^2}\right) E_c^o - 2\frac{a}{c^2} (\mathcal{L}_a - \mathcal{L}_c) \sum_{n=1}^{\infty} C_n J_1\left(x_n \frac{a}{c}\right).$$
(39)

Electrode Potential

The electrostatic potential P(r, z) at any point in the electrolyte is given by Eq. (9):

$$P(r, z) = C_0 + \sum_{n=1}^{\infty} C_n J_0 \left(x_n \frac{r}{c} \right) e^{-x_n z/c}.$$
 (9)

Use of Eq. (39) for C_0 in the above gives

$$P(r, z) = V' - \left(\frac{a}{c}\right)^2 E_a^o - \left(\frac{c^2 - a^2}{c^2}\right) E_c^o - \frac{2a}{c^2} (\mathcal{L}_a - \mathcal{L}_c) \sum_{n=1}^{\infty} C_n J_1 \left(x_n \frac{a}{c}\right) + \sum_{n=1}^{\infty} C_n J_0 \left(x_n \frac{r}{c}\right) e^{-x_n z/c}.$$
(40)

The relationship between the electrostatic potential P(r, 0) in the electrolyte near the metal surface and the electrode potential E(r, 0) vs a standard reference electrode is given by Eq. (10). Use of Eq. (40) with z = 0 in Eq. (10) gives

$$E(r,0) = \left(\frac{a}{c}\right)^{2} E_{a}^{o} + \left(\frac{c^{2}-a^{2}}{c^{2}}\right) E_{c}^{o} + \frac{2a}{c^{2}} \left(\pounds_{a} - \pounds_{c}\right) \sum_{n=1}^{\infty} C_{n} J_{1}\left(x_{n} \frac{a}{c}\right) - \sum_{n=1}^{\infty} C_{n} J_{0}\left(x_{n} \frac{r}{c}\right),$$
(41)

where the C_n are determined from Eq. (35). Again the indeterminate constant V' vanishes in the final expression.

Local Current Density

The local current density i(r, 0) along the metal surface is related to the electrostatic potential P(r, z) by Eq. (5). Performing the differentiation on P(r, z) as given in Eq. (40) and inserting the result in Eq. (5) yields

$$\frac{i(r,0)}{\sigma} = \frac{1}{c} \sum_{n=1}^{\infty} C_n x_n J_0\left(x_n \frac{r}{c}\right), \tag{42}$$

where again the set of C_n is determined from Eq. (35).

Total Anodic Current

The total anodic current is related to the local current density by [1, 2, 8]:

$$I_{\text{anodic}} = \int_{r=0}^{a} \int_{\theta=0}^{2\pi} i(r,0) r \, dr \, d\theta \tag{43}$$

or

$$I_{\text{anodic}} = 2\pi \int_{r=0}^{a} i(r, 0) \, r \, dr \tag{44}$$

Use of Eq. (42) in (44) gives

$$I_{\text{anodic}} = \frac{2\pi\sigma}{c} \int_{r=0}^{a} \sum_{n=1}^{\infty} C_n x_n J_0\left(x_n \frac{r}{c}\right) r \, dr \tag{45}$$

or

$$I_{\text{anodic}} = \frac{2\pi\sigma}{c} \sum_{n=1}^{\infty} \int_{r=0}^{a} C_n x_n J_0\left(x_n \frac{r}{c}\right) r \, dr.$$
(46)

The integral can be evaluated from Eq. (19), so that the result, omitting a few steps, is

$$\frac{I_{\text{anodic}}}{\sigma} = 2\pi a \sum_{n=1}^{\infty} C_n J_1\left(x_n \frac{a}{c}\right), \tag{47}$$

where again the set of C_n is determined from Eq. (35).

MATHEMATICAL ANALYSIS FOR A THIN-LAYER ELECTROLYTE

If the electrolyte is a thin layer of height b instead of bulk liquid, the boundary condition given by Eq. (8) is replaced by the requirement that there is no current flow across the outer boundary of the electrolyte:

$$\left[\frac{\partial P(r,z)}{\partial z}\right]_{z=b} = 0.$$
(48)

The other boundary conditions are the same as for the bulk case. The general solution of Laplace's equation subject to the restrictions of Eqs. (6), (7), and (48) is

$$P(r, z) = C_0 + \sum_{n=1}^{\infty} C_n \cosh\left[\frac{x_n}{c} (b-z)\right] J_0\left(x_n \frac{r}{c}\right).$$
(49)

The coefficients C_0 and C_n are evaluated from Eqs. (15a) and (15b), as was done for the case of bulk electrolyte. The procedure is exactly the same as for the bulk case; hence only the results are listed below.

The coefficients C_n are given by the systems of simultaneous equations

$$C_{m}\left\{\left[1+\mathcal{L}_{c}\frac{x_{m}}{c}\tanh\left(x_{m}\frac{b}{c}\right)\right]\frac{J_{0}^{2}(x_{m})}{2}+\left(\mathcal{L}_{a}-\mathcal{L}_{c}\right)\frac{x_{m}}{c}\left(\frac{a}{c}\right)^{2}\tanh\left(x_{m}\frac{b}{c}\right)A_{m}\right\}\right.$$
$$\left.+\frac{a}{c^{2}}\frac{\left(\mathcal{L}_{a}-\mathcal{L}_{c}\right)}{\cosh\left(x_{m}\frac{b}{c}\right)}\sum_{\substack{n=1\\n\neq m}}^{\infty}C_{n}W_{nm}\sinh\left(x_{n}\frac{b}{c}\right)=-\frac{1}{\cosh\left(x_{m}\frac{b}{c}\right)}\frac{\left(E_{a}^{o}-E_{c}^{o}\right)}{x_{m}}\left(\frac{a}{c}\right)J_{1}\left(x_{m}\frac{a}{c}\right),$$
(50)

where again A_m and W_{nm} are defined by Eqs. (35a) and (35b) respectively and m takes on the values 1 through k successively.

The constant C_0 is given by

$$C_{0} = V' - \left(\frac{a}{c}\right)^{2} E_{a}^{0} - \left(\frac{c^{2} - a^{2}}{c^{2}}\right) E_{c}^{0} - \frac{2a}{c^{2}} \left(\pounds_{a} - \pounds_{c}\right) \sum_{n=1}^{\infty} C_{n} \sinh\left(x_{n} \frac{b}{c}\right) J_{1}\left(x_{n} \frac{a}{c}\right),$$
(51)

$$E(r, 0) = \left(\frac{a}{c}\right)^{2} E_{a}^{o} + \left(\frac{c^{2}-a^{2}}{c^{2}}\right) E_{c}^{o} + \frac{2a}{c^{2}} \left(\pounds_{a} - \pounds_{c}\right) \sum_{n=1}^{\infty} C_{n} \sinh\left(x_{n}\frac{b}{c}\right) J_{1}\left(x_{n}\frac{a}{c}\right)$$
$$- \sum_{n=1}^{\infty} C_{n} \cosh\left(x_{n}\frac{b}{c}\right) J_{0}\left(x_{n}\frac{r}{c}\right).$$
(52)

The local current density is given by

$$\frac{i(r,0)}{\sigma} = \frac{1}{c} \sum_{n=1}^{\infty} C_n x_n \sinh\left(x_n \frac{b}{c}\right) J_0\left(x_n \frac{r}{c}\right),$$
(53)

and the total anodic current is

$$\frac{I_{\text{anodic}}}{\sigma} = 2\pi a \sum_{n=1}^{\infty} C_n \sinh\left(x_n \frac{b}{c}\right) J_1\left(x_n \frac{a}{c}\right).$$
(54)

When $b \to \infty$, C_n (thin-layer) \times sinh $(x_n b/c) \to C_n$ (bulk), so that the expressions for thinlayer electrolyte reduce to the corresponding equations for a bulk electrolyte for large b.

The various expressions for the thin-layer and bulk cases are summarized in Table 1.

PREVIOUS CASE OF EQUAL POLARIZATION PARAMETERS

When $\mathcal{L}_a = \mathcal{L}_c = \mathcal{L}$, the preceding results for bulk and thin-layer electrolytes reduce to

$$E(r, 0) = \left(\frac{a}{c}\right)^{2} E_{a}^{o} + \left(\frac{c^{2} - a^{2}}{c^{2}}\right) E_{c}^{o} + 2\left(\frac{a}{c}\right) (E_{a}^{o} - E_{c}^{o}) \sum_{n=1}^{\infty} \frac{J_{1}\left(x_{n} \frac{a}{c}\right)}{x_{n}\left[1 + \pounds \frac{x_{n}}{c} Q\right] J_{0}^{2}(x_{n})} J_{0}\left(x_{n} \frac{r}{c}\right),$$
(55)

and

Table 1—Summary of Relationships for Coplanar Circular Electrodes under a Bulk and a Thin-Layer Electrolyte. The hyperbolic functions in the upper part of the brackets apply to thin layers, and are replaced by 1 for a bulk electrolyte. The numbers in parentheses refer to equation numbers in the text.

$$Coefficients C_n$$

$$C_m \left[\left[1 + L_c \frac{x_m}{c} \left\{ \frac{\tanh\left(x_m \frac{b}{c}\right)}{\text{or}} \right\} \right] \frac{J_0^2(x_m)}{2}$$

$$+ (L_a - L_c) \frac{x_m}{c} \left(\frac{a}{c}\right)^2 \left[\frac{J_0^2\left(x_m \frac{a}{c}\right) + J_1^2\left(x_m \frac{a}{c}\right)}{2} \right] \left\{ \frac{\tanh\left(x_m \frac{b}{c}\right)}{\text{or}} \right\} \right]$$

$$+ \frac{a}{c^2} \left\{ \frac{L_a - L_c}{\cosh\left(x_m \frac{b}{c}\right)} \sum_{\substack{n=1\\ n \neq m}}^{\infty} C_n \left\{ \frac{\sinh\left(x_n \frac{b}{c}\right)}{\text{or}} \right\} + \frac{x_n^2}{x_n^2 - x_m^2} \left[J_1\left(x_n \frac{a}{c}\right) J_0\left(x_m \frac{a}{c}\right) \right] \right]$$

$$- \frac{x_m}{x_n} J_0\left(x_n \frac{a}{c}\right) J_1\left(x_m \frac{a}{c}\right) \right] = - \frac{1}{\left\{ \frac{\cosh\left(x_m \frac{b}{c}\right)}{\text{or}} \right\}^{\frac{n}{2}} \frac{x_n^2 - x_m^2}{2} \left[J_1\left(x_n \frac{a}{c}\right) J_0\left(x_m \frac{a}{c}\right) \right] \right\}$$
Electrode Potential
$$E(r, 0) = \left(\frac{a}{c}\right)^2 E_a^n + \left(\frac{c^2 - a^2}{c^2}\right) E_c^n + \frac{2a}{c^2} (L_a - L_c) \sum_{n=1}^{\infty} C_n \left\{ \frac{\sinh\left(x_n \frac{b}{c}\right)}{\text{or}} \right\} J_1\left(x_n \frac{a}{c}\right)$$

$$- \sum_{n=1}^{\infty} C_n \left\{ \frac{\cosh\left(x_n \frac{b}{c}\right)}{\text{or}} \right\} J_0\left(x_n \frac{r}{c}\right). \quad (41, 52)$$
Local Current Density
$$\frac{i(r, 0)}{a} = \frac{1}{c} \sum_{n=1}^{\infty} C_n x_n \left\{ \frac{\sinh\left(x_n \frac{b}{c}\right)}{\text{or}} \right\} J_0\left(x_n \frac{r}{c}\right) \quad (42, 53)$$
Total Anodic Current
$$\frac{I_{anodic}}{a} = 2\pi a \sum_{n=1}^{\infty} C_n \left\{ \frac{\sinh\left(x_n \frac{b}{c}\right)}{\text{or}} \right\} J_1\left(x_n \frac{a}{c}\right)$$

$$(47, 54)$$

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(47, 54)

$$\frac{i(r,0)}{\sigma} = -2 \frac{a}{c^2} \left(E_a^{\rm o} - E_c^{\rm o} \right) \sum_{n=1}^{\infty} \frac{J_1\left(x_n \frac{a}{c}\right)Q}{\left[1 + \pounds \frac{x_n}{c} Q\right] J_0^2(x_n)} J_0\left(x_n \frac{r}{c}\right), \tag{56}$$

$$\frac{I_{\text{anodic}}}{\sigma} = -4\pi \frac{a^2}{c} \left(E_a^o - E_c^o \right) \sum_{n=1}^{\infty} \frac{J_1^2 \left(x_n \frac{a}{c} \right) Q}{x_n \left[1 + \pounds \frac{x_n}{c} Q \right] J_0^2(x_n)} , \qquad (57)$$

where in each of the above

$$Q = \begin{cases} 1, & \text{bulk electrolyte,} \\ \\ \tan h\left(x_n \frac{b}{c}\right), & \text{thin layer.} \end{cases}$$
(58)

These equations are the same as those previously reported except that the signs of electrode potentials in Eqs. (55) through (57) now conform to the convention that the noble direction is the more positive.

NUMERICAL EVALUATION

Values of \mathcal{L}_{a} and \mathcal{L}_{c}

In general the anode and cathode have different polarizabilities (the two electrode potentials respond differently to the passage of current). In many instances the anode is the less polarizable. This is illustrated by many electrode kinetic studies carried out under carefully controlled conditions. With iron, for example, in a variety of deaerated electrolytes, anodic Tafel slopes of 30 to 80 mV/decade have been observed, while the cathodic slopes were 120 mV/decade [31-34]. Other metals in the iron group (nickel and cobalt) have been observed to behave similarly [35].

To cite two more examples, cadmium undergoes self-dissolution to Cd^{+2} in acids by two consecutive single-electron transfer reactions, and indium goes to In^{+3} through three consecutive single-electron transfers. The observed anodic Tafel slopes are 40 to 50mV/decade (0°C) for cadmium [36] and 22 mV/decade for indium [37], in good correspondence with the theoretical values of 2.303 RT/(3/2)F and 2.303 RT/(5/2)F respectively. The hydrogen-evolution reaction on both surfaces gave cathodic Tafel slopes of 115 mV/decade and 120 mV/decade respectively, indicative of a single-electron transfer characterized by a theoretical value of 2.303 RT/(1/2)F.

In more practical situations where conditions are not as well defined, Tafel behavior is not always observed, but instead polarization curves sometimes display segments which are approximately linear in current (rather than the logarithm of current) over a considerable potential range, as discussed earlier. In these instances the cathode often again

and:

is the more polarizable. Figure 3a shows a schematic Evans diagram [38] for a bimetallic couple under cathodic control. Other possibilities, however, include anodic control (Fig. 3b) and mixed control (Fig. 3c). This last case would approximate earlier treatments [1, 2, 8] for $\mathcal{L}_a = \mathcal{L}_c$.

Values of Wagner linear polarization curves compiled [1, 2] from the literature indicate that \mathcal{L}_a is generally of the order of 1 to 10 cm while \mathcal{L}_c is usually 10 to 100 cm, although there are exceptions. In data tabulated by Gouda and Mourad [15] for steel in a variety of neutral to basic solutions both with and without added chloride, the cathodic slope |dE/di| varied from 1.5 times to approximately 20 times the anodic slope but with most ratios in the range of 5 to 10. Specific conductivities were listed for only three solutions, for which values of \mathcal{L}_a are calculated to be 0.9, 1.9, and 48 cm, with corresponding \mathcal{L}_c values of 7.5, 12.2, and 193 cm respectively.



Fig. 3—Schematic representations of cathodic, anodic, and mixed control with L_a fixed at 1 cm in each case

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Bulk Electrolyte

For the calculations in this section, the anodic polarization parameter is fixed at $\mathcal{L}_a = 1 \text{ cm}$. Figure 4 shows the potential distribution E(r, 0) for $\mathcal{L}_a = 1 \text{ cm}$ and $\mathcal{L}_c = 10 \text{ cm}$ for fixed values of $E_a^0 = 0.00 V$ and $E_c^0 = 1.00 V$. The coefficients C_n were calculated up to C_{100} using the system of simultaneous equations generated by Eq. (35). These simultaneous equations were solved using the CDC 3800 computer, and the coefficients were then substituted in Eq. (41) to obtain the electrode potential distribution. Convergence was assessed by numerical evaluation. The computer program is given in Appendix B.

Figure 4 also shows potential distribution plots for $\mathcal{L}_a = \mathcal{L}_c = 1$ cm and $\mathcal{L}_a = \mathcal{L}_c = 10$ cm, calculated from Eq. (55). It is evident that the potential behavior of the electrodes with unequal anodic and cathodic polarization parameters cannot be deduced from the two separate curves for the equal polarization parameters.

The corresponding current density curves for the three systems are shown in Fig. 5. It is seen that the values for the case of unequal parameters are intermediate between the two cases where $\mathcal{L}_a = \mathcal{L}_c = 1$ cm and $\mathcal{L}_a = \mathcal{L}_c \approx 10$ cm.



Fig. 4—Comparison of electrode potential distributions for equal and unequal polarization parameters with bulk electrolyte (anode radius a = 0.5 cm, cathode radius c = 1.0cm, $E_a^o = 0.00$ V, and $E_c^o = 1.00$ V)



Fig. 5—Current distributions corresponding to the electrode potential distributions in Fig. 4

Figure 6 shows the potential distribution calculated from Eq. (41) with n = 100 for a fixed value of $\mathcal{L}_a = 1$ cm but with variable \mathcal{L}_c . Corresponding current distribution curves calculated from Eq. (42) are shown in Fig. 7. When $\mathcal{L}_a = 1$ cm and $\mathcal{L}_c = 0.1$ cm, the galvanic couple is under anodic control, as depicted in Fig. 3b, and the electrode potentials across the metal surface of both components are polarized up near the potential of the uncoupled cathode.

When $\mathcal{L}_c >> \mathcal{L}_a$, such as $\mathcal{L}_a = 1$ cm and $\mathcal{L}_c = 100$ cm, the system is under cathodic control, as illustrated schematically in Fig. 3a. For this case, Fig. 3a predicts that the electrode potential would approach the values of the open-circuit potential of the anode and the current would be much less than for the case of anodic control (for fixed \mathcal{L}_a). These expected trends are verified in the results of the numerical analysis shown in Figs. 6 and 7.



Fig. 6—Distribution of electrode potential across circular cells under bulk electrolyte with \mathcal{L}_a fixed at 1 cm, combined with various values of \mathcal{L}_c (anode radius a = 0.5 cm, cathode radius c = 1.0 cm, $E_a^o = 0.00$ V, and $E_c^o = 1.00$ V)

Results in Figs. 6 and 7 for $\mathcal{L}_a = \mathcal{L}_c = 1$ cm also provide a check on the consistency of the present method with the previous relationships for equal polarization parameters. Both current and potential distributions calculated from the set of C_n resulting from Eq. (35) with \mathcal{L}_a and \mathcal{L}_c both equal to 1 cm agree with the results obtained from Eqs. (55) and (56).

One additional trend can be seen in Figs. 6 and 7. For this system of fixed \mathcal{L}_a , the more polarizable the cathode (the larger \mathcal{L}_c), the more uniform the potential and current distribution.

The total anodic current was calculated from Eq. (47) for the systems with \mathcal{L}_a fixed at 1 cm with variable \mathcal{L}_c . Results are listed in Table 2.

The total anodic current can also be calculated from the schematic Evans diagrams shown in Fig. 3. For the anodic branch

$$\left|\frac{dE}{dI}\right|_{a} = \frac{E_{\rm corr} - E_{a}^{o}}{I_{\rm corr}}$$
(59)





Fig. 7—Distribution of current across circular cells under bulk electrolyte with \mathcal{L}_a fixed at 1 cm and combined with various values of \mathcal{L}_c (anode radius a = 0.5 cm, cathode radius c = 1.0 cm, E_a^o = 0.00 V, and E_c^o = 1.00 V)

and for the cathodic branch

$$\left|\frac{dE}{dI}\right|_{c} = -\frac{E_{\rm corr} - E_{\rm c}^{\rm o}}{I_{\rm corr}}, \qquad (60)$$

where I_{corr} is the corrosion current (the total anodic current referred to earlier as I_{anodic}). $I_a = i_a A_a$ and $I_c = i_c A_c$, where A_a and A_c are the area of anode and cathode respectively. Use of Eqs. (1) and (2) in Eqs. (59) and (60) gives

<i>L_a</i> (cm)	<i>L_c</i> (cm)	$I_{\rm corr}/\sigma$, Calculated from Evans Diagrams: Eq. (61)	I_{anodic}/σ , Calculated from Eqs. (35) and (47)
1	0.1	0.760	0.607
	1	0.589	0.485
	2	0.471	0.401
	10	0.181	0.169
	20	0.103	0.099
	50	0.0445	0.0437
	100	0.0228	0.0227
10	10	0.0589	0.0576
	100	0.0181	0.0179
100	100	0.00589	0.00589

Table 2—Comparison of Total Current Calculated from Evans Diagrams and From Summation of Current Distribution Curves for Circular Couples Under Bulk Electrolyte (anode radius a = 0.5 cm, cathode radius c = 1.0 cm, $E_a^o = 0.00$ V, and $E_c^o = 1.00$ V)

$$\frac{I_{\text{corr}}}{\sigma} = \frac{E_c^o - E_a^o}{\frac{L_a}{A_a} + \frac{L_c}{A_c}},$$
(61)

where again I_{corr} has the same meaning of I_{anodic} in Eq. (47).

Values of I_{anodic}/σ calculated from Eq. (47) are also listed in Table 2. These calculated values agree with the results from the computer analysis for $\mathcal{L}_a = 1$ cm coupled with cathodic values of $\mathcal{L}_c = 50$ cm and $\mathcal{L}_c = 100$ cm, where the current distribution is uniform, as shown in Fig. 7. There is disagreement between the results of the Evansdiagram analysis and the computer analysis for those systems where there is a nonuniform distribution of current, and this divergence is greater the more pronounced the localized attack at the anode/cathode juncture.

Results for $\mathcal{L}_a = \mathcal{L}_c = 10$ cm and $\mathcal{L}_a = \mathcal{L}_c = 100$ cm are also included in Table 2. Current distribution plots published in an earlier report [1] were nearly uniform for the former system and exactly so for the latter. The current distribution for $\mathcal{L}_a = 10$ cm and $\mathcal{L}_c = 100$ cm was calculated from Eq. (42) and was also observed to be uniform (The plot is not shown here.) Thus there is good agreement between Evans-diagram analyses and computer calculations for the cases where there is a uniform current distribution.

Thus the classic Evans polarization diagrams cannot be used to accurately predict the value of galvanic currents unless the anode and cathode components each behave uniformly.

Thin-Layer Electrolyte

Figure 8 shows the electrode potential E(r, 0) for $\mathcal{L}_a = 1$ cm and $\mathcal{L}_c = 10$ cm for different electrolyte thicknesses. The coefficients C_n were computed up to C_{100} from Eq. (50) and were used in Eq. (52) to obtain the potential distributions. The computer program for thin layers is given in Appendix C.

Figure 8 shows that the potential distribution is almost uniform for bulk electrolyte, but with thin layers most of the polarization takes place near the anode/cathode juncture. The anode center and cathode outer edge are virtually unaffected by the presence of each other for the thinnest electrolyte of 0.001 cm.

The corresponding current distributions are shown in Fig. 9. The local current densities were calculated from Eqs. (50) and (53) with n = 100, except near r = 0.0 and r = 0.5, where 125 terms were used. For the thin layers there is a geometry effect in which the corrosion attack is concentrated near the anode/cathode boundary.



Fig. 8—Distribution of electrode potential for $\mathcal{L}_a = 1$ cm and $\mathcal{L}_c = 10$ cm for different electrolyte thicknesses (anode radius a = 0.5 cm, cathode radius c = 1.0 cm, $E_a^o = 0.00$ V, and $E_c^o = 1.00$ V)



Fig. 9—Current distribution for $\mathcal{L}_a = 1 \text{ cm}$ and $\mathcal{L}_c = 10 \text{ cm}$ for different electrolyte thicknesses. The cell parameters are the same as in Fig. 8. (Limits between which the local current densities oscillate as computed from Eq. (53) are indicated for the anodic points. Limits are not shown for the cathode but are approximately half the range of the anodic points)

Figure 10 shows the total anodic current calculated from Eq. (54) for two different combinations of \mathcal{L}_a and \mathcal{L}_c . In both cases the total current approaches values for the bulk for electrolyte thicknesses of approximately 0.1 to 0.3 cm.

Figure 11 compares the potential distribution for $\mathcal{L}_a = 1$ cm and $\mathcal{L}_c = 10$ cm to the cases of equal polarization parameters: $\mathcal{L}_a = \mathcal{L}_c = 1$ cm and $\mathcal{L}_a = \mathcal{L}_e = 10$ cm for an electrolyte thickness of 0.001 cm. Figure 12 shows a similar curve for $\mathcal{L}_a = 10$ cm and $\mathcal{L}_c = 100$ cm. In each case the potential distribution for the system of unequal parameters is not related in a simple manner to the individual distribution curves for each of the two equal parameters.



Fig. 10—Total anodic current computed as a function of electrolyte thickness for two different combinations of \mathcal{L}_a and \mathcal{L}_c (anode radius a = 0.5 cm, cathode radius c = 1.0 cm, $E_a^o = 0.00$ V, and $E_c^o = 1.00$ V)



Fig. 11—Comparison of electrode potential distribution for equal and unequal polarization parameters for a thinlayer electrolyte of thickness b = 0.001 cm (anode radius a = 0.5 cm, cathode radius c = 1.0 cm, E_a^o = 0.00 V, and E_c^o = 1.00 V)



Fig. 12—A second comparison of electrode potential distribution for equal and unequal polarization parameters for a thin-layer electrolyte of thickness b = 0.001 cm (anode radius a = 0.5 cm, cathode radius c = 1.0 cm, E_a^o = 0.00 V, and E_c^o = 1.00 V)

SUMMARY

A mathematical model has been developed to describe the distribution of potential and current across circular corrosion cells having unequal anodic and cathodic linear polarization parameters. This analysis is applicable to systems of bimetallic galvanic couples or to systems with a localized geometry effect, as in pitting corrosion.

The potential distribution in a system having unequal anodic and cathodic polarization parameters is not related in a simple manner to the separate distribution curves for the two cases where the polarization parameters are equal.

For bulk electrolyte the value of the electrode potentials depends on whether the system is under anodic, cathodic, or mixed control. Current distribution is more uniform for the more polarizable combinations of electrodes. Thus the total corrosion current calculated from the Evans diagram is in error if the individual current distributions are not uniform.

In thin-layer electrolytes there is a geometry effect in which the electrode polarization and current flow is concentrated near the anode/cathode juncture. In a typical system the tendency toward bulk behavior occurs at about 0.1 to 0.3 cm (1000 to 3000 μ m).

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Appendix A

RELATIONSHIP BETWEEN THE ELECTRODE POTENTIAL AND THE ELECTROSTATIC POTENTIAL AT THE METAL/SOLUTION INTERFACE

As pointed out by Bockris and Reddy [A1], it is impossible to measure the electrode potential of a metal/solution interface without introducing additional extraneous interfaces during the measurement process. This is illustrated in Fig. A1, where the electrode potentials of the coplanar anode and cathode are to be measured versus the reference electrode.



Fig. A1—Method of measuring the electrode potentials of a coplanar anode (A) and cathode (C)

In circuit 1 the measured electrode potential of the anode E_a vs the reference electrode is related to the potential differences across the various interfaces by

$$[\phi_A - P(r, 0)] + [P(r, 0) - \phi_{\text{Ref}}] + (\phi_{\text{Ref}} - \phi_{M'_1}) = E_a,$$
(A1)

where ϕ_A is the electrostatic potential "just inside" the metal A and P(r, 0) is the electrostatic potential in the solution "just outside" the metal [A2]. Similarly ϕ_{Ref} and $\phi_{M'_1}$ refer respectively to the electrostatic potential just inside the solid-phase reference electrode and just inside the connecting wire.

According to Bockris and Reddy [A1] the potential difference across a nonpolarizable interface such as ϕ_{Ref} /solution is a constant, so that

$$\phi_A + P(r, 0) - \phi_{\text{Ref}} + (\phi_{\text{Ref}} - \phi_{M'_1}) = V', \qquad (A2)$$

where V' is a constant. Use of this definition of V' in Eq. (A1) gives

$$V' - P(r, 0) = E_a, \quad 0 \le r < a.$$
 (A3)

Similarly measurement of the electrode potential of the cathode E_c in circuit 2 gives

$$(\phi_A - \phi_C) + [\phi_C - P(r, 0)] + [P(r, 0) - \phi_{\text{ref}}] + (\phi_{\text{ref}} - \phi_{M_1'}) = E_c.$$
(A4)

The ϕ_C terms in Eq. (A4) cancel, so that

$$V' - P(r, 0) = E_c, \quad a < r \le c.$$
 (A5)

Thus

$$V' - P(r, 0) = \begin{cases} E_a, & 0 \le r \le a \\ E_c, & a < r \le c \end{cases}$$
(A6)

Or denoting the electrode potential along the metal surface as E(r, 0) gives

$$V' - P(r, 0) = E(r, 0), \tag{A7}$$

which is Eq. (10) in the main text

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Appendix B

COMPUTER PROGRAM FOR COPLANAR CONCENTRIC CIRCULAR ELECTRODES WITH UNEQUAL POLARIZATION PARAMETERS UNDER BULK ELECTROLYTE

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15	F@RMAT (5%,*M*,9%,*%(M)*,13%,*F*,14%,*Y(M)*,11%,*B(M,1)*,10%,*B(M 10)*,9%,*B(M,50)*,8%,*B(M,100)*,//)
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	T3=PESJ(X(M),0)
	14=(13++2)/2.0
	15=12+14
	T6=X(M)+A/C
	17=FESJ(T0,C)
	18=BESJ(16,1)
	19=(17++2)+(18++2)
	T1C=((LA-LC)+X(M)+((A/C)++2)+T9)/(2,0+C)
	F=T5+T10
19	Y(M) = -(EA - EC) + (A/C) + (TB/X(M))
	N=1
20	1F (N.EG,M) 21.22
21	E(M,N)=P
	GO TE 30
22	112=x(M)/x(N)
	13=1.0/(1,0-(112++2))
	114=x(N)+A/C
	115=8ESJ(114,1)
	116=EESJ(114,0)
	117 = (115 + 17) - (112 + 116 + 18)
	E(P,N) = (LA - LC) + (A/(C + 2)) + 13 + 17
	Ge 76 30
50	N=N+1
	1F (N.LE.K) 20,31
51	FRINI 32, F, X(F), F, Y(F), E(F, 1), E(F, 10), B(F, 50), 3(M, 100)
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  812 READ 813, RH
  813 FORMAT (F10.0)
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  820 FRINT 83, RR
   83 FORMAT (////, 3x, 3HRR= F5,3)
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84 FORMAT C
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   FRINT 85
85 FREMAT (60X, +0+, 14X, 9+ +EESSEL2=)
   PRINT 86
86 FORMAT (5x,+N+,5%,+X(N)+,11x,+(SLPN+,12x,+XNRC+,10x,+RESSEL2+,9x,+
      1TERMILOCO, 9X, +ILCCALO,//)
       ILCCAL:0,0
       LO 89 N=1.K
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94 FOHMAT (////,Sx.3HRR= F5,3)
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96 FERMAT (60x,+U+,13x,14HCSLBN+BESSEL2=,3x,9HTERMPOTL=,7x,12H-CO-SLM
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15	LENTINUE		00003400
	RETURN		00003900
e	LO 10 I=1.NO	이 방법에 방법에 관련했다. 정말 것 같은 것 같은 것 같은 것 같이 많이	00004000
	IF(I,GT,9)60 10 2	C	00014101
	2ERE(1)=XJA5(1)		00004200
10	CONTINUE		00004300
	RETURN		00004400
50	EETA=(P1/4,0)+(2,1	0 • J&RE • 4, 0 • I • 1, 6)	00004500
	h1=8ETA+8.0		00004600
	w2=w1•w1		00004700
	LEHE(I)=BEIA-(HEL	D-1.C)/w1*(1;0+1.0/w2*(4.0*(7,0*HOLD=31,0)/3.0	00004800
	1 +1.0/~2.032,	0 • (83, C • FOLD • FELE • 982, U • HOLD • 5779, 0)/15, 0	00004900
	2 +1.0/~2.64.	0•/6949,C*F0LC*F7LC*F0LL-153855,U*H0LD*H0LD	0.0005000
	3 +1585743.0+1	GLD-6277237.01/105.01)))	07005100
	60 16 (11,12,13,1	4,15,16)_0RD+1	00005200
10	ENL		00005300
	FUNCTION BESULX.N)	BESJ 1
	LATA (RO= , 20278449	47E8),(H1=+852659891E/),(R2#.38831312263E6),	BESJ 2
	1(83=90578674277	E4), (H4=,108306963F3), (H5=-,73485335935),	BESJ 3
	2(H6=,29212672487E	-2),(R7=-,65050170571E-5),(RA=.64538018051F+A),	BESJ 4
	3150=,282784494788),(51=,21095247743E6),(52=,70046825147E3),	BESJ 5
	4(A0=2.5323420902E:	2), (A1=4, 2217704118E1), (A2=5, 2443314672E-1),	BFSJ 6
	5(80=,44884594896E.	3),(F1=,753220485/9E2),	B-SJ 7
	6(00=-1,23.59445551	E1),(C1=-2.7788921059),(C2=-4,9517399126E-2),	BESJ A
	7(01=,410055452352), (F=64,), (G=4,72236648221),	BESJ 9
	8110=,17496878239E	3),	845J 10
	A(RH0=.9808/274959	E7), (HE1=11425325721E/), (RR2=,40945213625E5),	BESJ 11
	91863= 6666 111985	6F3). (FH4=.57575414035F1). (PH2=27914475519F-1).	BESJ 12

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C(RR6=,73493132111E-4),(RR7=-,84306821641E-7), BESJ 13 C(SS0=,1961/454991E8),(SS1=,16711673184E6),(SS2=.60777258247E3), BESJ 14 F(FE0=,62836856631E3),(8F1=,97300094628E2), BESJ 15 F(FU0=,21185478331F3),(UE1=,46917127629E2), G(AA0=3,5451899975E2),(AA1=5,5544843021E1),(AA2=6,5223084285E=1), BESJ 16 BESJ 17 +(CC0=4,4822348228E1),(CC1=9,7348068764),(CC2=1,7725579145F-1) BESJ 18 BESJ 19 L=X+X BESJ 21 1F (N, EU, 0) GO TE 6 \$ 1F (N, EG. 1) GE TE 7 \$ GO TO 8 BESJ 15(0-5)1,1,2 21 6 1 P=((((R8+J+R7)+C+R6)+C+R5)+C+R4)+C \$ P=(((P+R3)+D+R2)+D+R1)+C+R0 BESJ 22 23 HESU = P/(((D+S2)+C+S1)+C+S0) & RETURN BESJ BESJ 24 2 IF(D,GT,G) GO TE 9 A=ABS(X) % Dat/C BESJ 25 F=((A2+D+A1)+D+A0)/((C+E1)+C+E0) BESJ 24 G= ((C2+D+C1)+C+C0)/(A+((D+D1)+C+L0)) BESJ 27 BESJ EESU = (COS(A) + (P+C)+SIN(A) + (P-C))/SCRT(A) 28 29 RETURN BESJ BESJ 30 IF(C-F)11.11,21 7 BESJ 11 F=((((((HH7*D*RH6)*D+FH5)*D+RH4)*C+FH3)*D+RR2)*D+RR1)*D+RR0 31 BESJ=X+P/(((D+SS2)+D+S51)+U+SSC) & HETURN BESJ 32 21 IF(C,GT,G) GO TE 9 BESJ 33 A=ABS(x) > D=F/C BESJ 34 BESJ 35 F=((AA2+D+AA1)+C+AA0)/((D+BE1)+F+HE0) G=((CC2+D+CC1)+F+CC0)/(A+((E+FL1)+L+LE0)) BESJ 34 BESJ 37 A=(C@S(A)+(U-P)+SIN(A)+(C+P))/SGR1(A) % 1F(X, 1, U)A=-A BESJ 34 EESJEA BESJ 39 RETURN 40 E FRINT 81, V BESJ 81 FORMAT(//15X+ERRER IN BES., N =+15) BESJ 41 GA TE 100 BESJ 42 FRINTOLIX BESJ 43 9 91 FFRMAT(//15X+ERRER IN BESLI ARGLMENT & TER LARGE, X = +E17.10) BESJ 44 100 BESJ=1.8300 BESJ 45 BESJ END 46 SUBREUTINE MATALC(A, X, NR, NV, ICE, LET, NACT) LIMENSION A(NACT, NACT), X(NACT, NACT) 000 001 IF(IL@) 1.2.1 002 003 1 LO 3 1=1,NH 004 LA 4 J=1,NR 4 x(1,.)=0,0 005 3 x(1,1)=1,0 000 NV=NA 007 2 LFT=1.0 008 009 NR1=NR+1 EA 5 K=1, NR1 010 1P1=K+1 011 F1VET=0.0 012 LP & Isk, NH 013 014 L=ABSF(A(1,K)) 015 1F(Z-PIVOT) 6,6,7

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,	£1461	016
'	1961-2 196-1	017
	CONTINUE	015
c		C19
		020
		021
		022
	IFTIFREN) 1JJ11.10	023
10		024
		025
	A(LPH, J) =A(N, J)	026
14	A(K, u)=2	027
	Le 13 J=1,NV	028
	2=x(1PR,J)	020
	X([PR,J)=X(K,J)	010
13	X(K, J)=2	0.50
	LET==DET	031
11	LET=LET+A(K,K)	032
	PIVET=1.G/A(K.K)	033
	L0 14 J=141.NR	034
	A(K, J) = A(K, J) + PIVET	035
	CO 14 [=[H1,NH	036
14	A(1,) = A(1, J) - A(1, K) + A(K,)	037
	LA 5 JEL,NV	038
	1F(X(K,J)) 15,5,15	039
15	X(K,_)=X(K,J)=PIVET	040
	L0 16 [=[R1,NR	041
16	X(1,_)=X(1,J)=A(1,K)=X(K,_)	042
5	CONTINUE	043
	[F(A(NR, NH)) 17, 5, 17	044
17	LET=LET +A(NR,NR)	045
	FIVET=1,0/A(NR,NE)	046
	L0 18 J=1,NV	047
	X(NR, J)=X(NR, J)+F1V01	048
	L0 16 K=1,NR1	049
	IshR-K	050
	SUF=0.0	051
	EG 19 L=1, NR1	052
19	SUF=SUM+A(1,L+1)+X(L+1,_)	053
1.4	X(1,_)=X(1,J)=SLM	054
		055

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Appendix C

COMPUTER PROGRAM FOR COPLANAR CONCENTRIC CIRCULAR ELECTRODES WITH UNEQUAL POLARIZATION PARAMETERS UNDER THIN-LAYER ELECTROLYTE

FREGRAM UNEGTHIN

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THIS PROGRAM COMFLITES (1) CLERENT LISTRIBLIIEN (2) PETENTIAL DISTRIBUTION (3) TETAL ANGEIC CURHENT FOR CONCENTRIC CIRCULAR ELECTRECES COVERED BY THIN-LAYER FLECTROLYTE FOR THE CASE WHERE ANEDIC AND CATHEDIC WAGNER PULARIZATION FARAMETERS ARE NET FOLAL. A=RALIUS OF ANOLE C=RALIUS OF CATHEDE RECISTANCE ALONG RADILS ELENGTH=THICKNESS OF ELECTHELYTE LAYER ARER/C LASANODIC MAGNER POLAFIZATION FARAMETER LC=CATHODIC WAGNER PELAFIZATIEN FAMAMETER X(M)=MTH ZERO OF RESSEL FUNCTION OF GRUEN JORD P(R,0)=INTERFACIAL POTENTIAL ALONG THE METAL SURFACE E(R, 0) = POTENTL= ELECTECE PETENTIAL ALENG THE METAL SURFACE E(R,0)=CONSTANT-F(R,0), WITH THE CENSTANT CANCELLING OUT IN THE FINAL EXPRESSION, SO CENSTANT IN EFFECT CAN BE SET EQUAL TO LERG. THE COEFFICIENTS CO AND CSUEN ARE DEFINED IN THE FOLLOWING EQUATION F(H, 0) = C + SUP (C = (X(N) + F/C))0 N=1 N 0 ITETAL #TOTAL ANELIC CURRENT DIVIDEL BY THE CONDUCTIVITY ILECAL=LOCAL CURRENT DENSITY DIVIDED BY THE CONDUCTIVITY REAL LA, LC, ITOTAL, ILCCAL LIPEASIEN X(100),Y(100,100),H(100,100) F1=3,1415926536 #=100 1 REAC 10, A, C, BLENGTH, LA, LC, EA, FL 10 FORMAT (7F13.0) 1A: F10, 5, 5x, 3HLC: F10.5, 5x, 3HEA: F10.5, 5x, 3HEC: F10.5, ////) FRINT 12 12 FORMAT (1X, GENERATION OF THE SYSTEM OF SIMULTANEOUS EQUATIONS USF 11 TO SOLVE FOR THE COEFFICIENTS CO AND CSUBN+,///) FRINT 13 13 FRAMAT ((1X, NOT ALL THE CREFFICIENTS SO GINERATED ARE LISTED RELEN 1.,//) FRINT 14 14 FORMAT ((1X, +AS & PARTIAL CHECK, WHEN NEM, THE VALUE OF B(M, N) IS " 11VEN HY P.,////)

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FRINT 15
15 FORMAT (5X,+M+,5X,+X(F)+,13X,+F+,14X,+Y(F)+,11X,+B(M,1)+,10X,+B(M,
      110) . 9x, . H(M, 50) . 8x, . E(M, 100) . //)
С
       THIS PART OF THE PROGRAM GENERATES THE SYSTEM OF SIMULTANEOUS
С
C
       EGUATIONS USED IN SOLVING FOR THE COEFFICIENTS CO AND CSUBN.
       .CRE=1
       NO=K
       CALL HESZERA (JERE, NE, X)
       LO 4L M=1.X
XMBC=X(M)+BLENGTH/C
       SINHXMBC=(EXP(XMEC)-EXP(-XMEC))/2.0
       CESHXMHC=(EXP(XMEC)+EXP(-XMPC))/2.0
       TANHXMBC=SINHXMHC/COSFXMBC
       11=X(M)/C
       12=1,0+(LC+T1+TANHXMEC)
       13=865J(X(M),0)
       14=(13++2)/2.0
       15=12+14
       16=X(M)+A/C
       17=865J(16,0)
       18=FESJ(16,1)
19=(17++2)+(18++2)
       110=((LA-LC) + X(M) + ((A/C) + + 2) + TS + TANH XMBC)/(2,0+C)
       F=15+110
   19 Y(M)=-(1,0/COSHXMFC) + (EA-EC) + (A/C) + (18/X(M))
   N=1
20 IF (N.EC.M) 21.22
21 E(M.N)=P
G0 TE 30
    22 112=X(M)/X(N)
       113=1.0/(1, J-(112++2))
       114=X(N) .A/C
       115=6ESJ(114,1)
116=6ESJ(114,0)
       117=(115+17)-(112+116+18)
       XNEC=X(N) + BLENGTF/C
       SINHXNBC=(EXP(XNEC)-EXP(-XNEC))/2.0
       E(M, N)=(LA-LC)+(A/(C++2))+(1.C/CESFXMFC)+T13+T17+51NHXNBC
       GO TE 30
   30 N=N+1
   IF (N.LE,K) 20,31
31 PRINT 32,M,X(M),F,Y(M),E(M,1),B(M,10),B(M,50),3(M,100)
32 F0HMAT (3X,14,7(3X,E13.5))
    40 CONTINUE
C
       SELUTION OF THE SYSTEM OF & SIMULTANEEUS FQUATIONS
C
C
```

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THE SUBROUTINE REPLACES THE CONSTANT VECTORS WITH THE SOLUTION VECTORS, THUS, THE CONSTANT VECTORS Y(M) DEFINED IN STATEMENT 19 ARE REPLACED BY THE SOLUTION VECTORS. THESE SOLUTION VECTORS ARE LATER RELABELLED AS COURN. C C C С С CALL MATALG(8, Y, 100, 100, 0, DET, 100) С CALCULATION OF THE CEEFFICIENT CO С C FRINT 41 41 FORMAT (////,1H1,+CALCULATION OF THE COEFFICIENT CO+,//) FRINT 42 42 FORMAT (20x, 0450LLTIGN, 23x, 124. (X(N) • A/C), 2X, 1045INH(X(N) •, 7X, 134 1Y(N) • 5INHXNHC, 3X, 14HRLNAING SUM EF) FRINT 43 43 FORMAT (28x,7HVECTOR5,9x,8HX(A)+A/C,8x,+1+,17x,10HBLENGTH/C),2X,9H 1.PESSEL1:, /X, 9HTERMZERG:) FRINT 44 44 FERMAT (5X,+N+,9X,+X(N)+,11X,+Y(N)+,12X,+XNAC+,11X,+BESSEL1+,9X,+S 11N+XNHC ., 7X, . TERM7ER6 ., 10X, . 276TAL .//) ZTETAL=0.0 LP 46 N=1.K XNAC=X('.) • A/C EESSEL1=BESJ(XNAC,1) XNEC=X(1.)+BLENGTF/C SINHXNHC=(EXP(XNEC)-EXF(=XNEC))/2.0 TEHMZERESY(N) .BESSEL1.SINFXNBC 2TETAL=ZTATAL+TERMZERE FRINT 45, N, X(N), Y(N), XNAC, BESSEL1, SINHXNBC, TERMZERO, ZTOTAL 45 FORMAT (3X, 14, 7(3X, E13, 5)) 46 CENTINUE 21=((A/C) ...2) .EA 22=(1.0-((A/C) +2))+EC 23=(A/(C++2))+(2,0+(LA-LC))+2767AL PRINT 47,21,22,23 47 FORMAT (////,5X,3H21= E13,5,5X,3H23= E13,5) 60=-11-12-13 FRINT 48,00 48 FERMAT (////,5x,3+00= E13,5) С CALCULATION OF THE TETAL ANEDIC CURRENT C С FRINT 70 70 FORMAT (////.1H1.+CALCULATION OF THE TOTAL ANODIC CURRENT+,//) FRINT 71 71 FERMAT (1X, +ITUTAL =TETAL ANEDIC CUMPENT DIVIDED BY THE CONDUCTIVIT 11 PHINT 72 72 FERMAT (45x, 6HX(N)+A/C, 6X, 12H_ (X(N)+A/C), 2X, 10HSINH(X(N)+, 7X, 14H2 1,0.P1.A.CSUHN)

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```
FRINT 73
75 FORMAT (60X.+1+.17X.10HELENGTH/C),2X,17H+5INHXVBC+BESSEL1)
     FRINT 74
 74 F@HMAT (5x, *N*, 5x, *X(N)*, 11x, *CSLEN*, 11x, *XNAC*, 11x, *BESSEL1*, 9x, *
15INHXNBC*, 7X, *#TERMITET*, 5X, *TIETAL*, //)
     ITETAL=0,0
     LO 79 N=1.K
     XNAC=X(N)=A/C
     EESSEL1=BESJ(XNAC.1)
     CSLPN=Y(N)
     XNBC=X(N) +BLENGTF/C
     SINHXNBC=(EXP(XNEC)-EXP(-XNEC))/2.0
     TERMITOT=2, 0+PI+A+CSLEN+BESSEL1+SINHXNBC
     ITETAL=ITOTAL + TEHMITET
 FRINT 75, N, X(N), (SUAN, XNAC, HESSEL1, SINHXNAC, TERMITOT, ITOTAL
75 FERMAT (3X, 14, 7(3X, E13, 5))
 79 CONTINUE
     CALCULATION OF THE LECAL CURRENT PENSITY
 FFINT BO
BO FERMAT (////,1H1,+CALCULATIEN OF THE LOCAL CURRENT DENSITY+,//)
 FRINT 61
R1 FOHMAT (1X, .ILUCAL=LECAL CURRENT DENSITY DIVIDED BY THE CONDUCTIVI
   114.///)
810 READ 811, ROUT
811 FORMAT (F10.0)
812 READ 813, RH
813 FORMAT (F10.0)
    IF (HR.NE.HOUT) E20.90
820 FRINT 83,RR
P3 FORMAT (////,3X,3HRR= F5,7)
 FRINT 84
84 FORMAT (45x, 6HX(N)+R/C, 6X, 12H_ (X(N)+F/C), 2X, 10HSINH(X(N)+, 7X, 16H)
   11/0) + CSLBN + X(N))
 FRINT 65
85 FORMAT (60X,+0+,17X,1CHELENGTH/C),2X,17H+SINHXNBC+BESSEL2)
     FRINT 86
 86 FORMAT (5x, +N+, 9X, +X(N)+, 11X, +CSLEN+, 12X, +XNRC+, 10X, + 3ESSEL2+, 9X, +
   1SINHXNBC.,7X, .= TERMILEC., SX, . ILECAL. //)
     ILECAL=0.0
     LC 89 N=1.K
     XAHC=X(A)+RR
     EESSEL2=BESJ(XNRC.0)
     (SLEN=Y(N)
     XNBC=X(N)=BLENGTF/C
     SINHXNBC:(EXP(XNEC)-EXP(=XNEC))/2.0
TERMILUC:(1.0/C)+CSUEN+X(N)+SINHXNEC+EESSEL2
     ILECAL = ILOCAL . TERMILEC
     FRINT 87, N, X(N), CSUBN, XNRC, EESSEL2, SINHXNPC, TERMILOC, ILOCAL
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87 FORMAT (3X, 14, 7(3X, E13.5))
   85 CONTINUE
       GO TE 812
С
       CALCULATION OF THE PETENTIAL DISTRIBUTION ALONG THE HETAL SURFACE
CC
C
   96 FRINT 91
91 FREMAT (////,1H1,+CALCULATION OF THE FOTENTIAL DISTRIBUTION+,//)
   FRINT 92
92 FORMAT (1X, PROTENTLEE(R, 0)=ELFCTREEE POTENTIAL ALONG THE METAL SUR
     1FACF .....
  920 HEAD 921,88
  921 FORMAT (F10.3)
       IF (FR, NE. RCUT) $30.100
  930 FRINT 94, RH
94 FORMAT (////, 3x, 3+RR= F5, 3)
   FRINT 95
95 FRHMAT (43x.12H, (X(N)*F/C),2%,1CHCESH(X(N)*,7X,13HCSJBN*RESSFL2,3
1X,14HRUNNING SUM RF)
      FRINT 96
   96 FREMAT (44x. +0+.17x.1CHELENGTH/C). JX. 10++COSHXVBC=.6X.9HTERMPATLE.
     16X,12H-CO-SUMPOTL=)
       PRINT 97
   97 FORMAT (5x, .N., 9x, .X(N) ., 11x, .CSLBN.., 10x, .BESSEL 2., 9x, .COSHXNGC., 8
      1X. . TERMFOIL . HX. . SUMFEIL . dx. . FEIENTL . //)
       SUMPETL=0.C
       LO 99 N=1.K
       XNHC=X(N)+RH
       BESSEL2=BESJ(XNAC.0)
       CSLEN=Y(N)
       XNBC=X(N)=BLENGTF/C
       COSHXNBC = (EXP(XNEC) + EXP(-XNEC))/2.0
       TERMFOTL CSUBN. HESSELZ.COSHXNFC
       SUMPETL=SUMPOTL+TERMPETL
       FETENTL=- (CJ+SUMFETL)
   FRINT 98, N, X(N), CSURN, BESSEL2, CESHXNBC, TERMPOTL, SUNPOTL, POTENTL
98 FORMAT (3X, 14, 7 (3X, E13, 5))
   99 CONTINUE
       G0 TE 920
  10C END
```

SUBROUTINES BESZERO (JORD, NO, ZERO), FUNCTION BESJ (X, N), AND MATALG (A, X, NR, NV, IDO, DET, NACT) ARE GIVEN IN APPENDIX B.

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