# Platinum-Catalyzed Oxidations of Organic Compounds by Ferric Sulfate: Use of a Redox Fuel Cell to Mediate Complete Oxidation of Ethylene Glycol by Dioxygen at 80°C

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oxidized under mild conditions  $(60-80^{\circ}C, 3 \text{ } M \text{ } H_2SO_4$  in water)<br>by Fe(III) using a catalytic amount of Pt black to generate<br>Fe(II) and protons. The extent of oxidation depended on the<br>structures of the organic re **by oxidation of ethylene glycol was reoxidized to Fe(III) at the** the Fe(III)/Pt system and complete oxidation of ethylene anode of the cell. A V(V)/V(IV)/HNO<sub>3</sub>/O<sub>2</sub> redox system was glycol to carbon dioxide under mild c anode of the cell. A  $V(V)/V(IV)/HNO<sub>3</sub>/O<sub>2</sub>$  redox system was **used to catalyze the electrochemical reduction of dioxygen.** gen as stoichiometric oxidant in a redox fuel cell operating **Both the anode and cathode consisted of WDF graphite felt**, at 80°C. This is the first report of a Both the anode and cathode consisted of WDF graphite felt, at 80°C. This is the first report of a redox fuel cell operating the surface of which had been oxidized by boiling concentrated with ethylene glycol as fuel (5, 1 **cm<sup>3</sup> of graphite felt was obtained at a cell voltage of 197 METHODS mV.**  $\circ$  1996 Academic Press, Inc.

electrochemically oxidized to Fe(III) at the anode, and O<sub>2</sub> directly or from Aldrich (as the pentanydrate) and recrys-<br>was reduced to H<sub>2</sub>O at the cathode using a V(IV)/V(V)/ tallized from 6 M H<sub>2</sub>SO<sub>4</sub> before use. Solut  $HNO<sub>3</sub>$  redox system developed by Kummer and Oei (4–6). The mately 0.5 *M* in ferric sulfate and 0.8 *M* in sulfuric acid 3 redox system developed by Kummer and Oei (4–6). (Aesar, puratronic grade) were prepared and [ A Nafion membrane separated the anode and the cathode<br>compartments of the call and allowed transfer of protons determined by titrating aliquots of the solution with EDTA compartments of the cell and allowed transfer of protons determined by titrating aliquots of the solution with EDTA<br>from the anolyte to the catholyte. Both the anode and the  $at 40^{\circ}$ C using N-(4-methoxyphenyl)-1,4-pheny from the anolyte to the catholyte. Both the anode and the anode and the hydrochloride (Aldrich) as the indicator. cathode consisted of WDF graphite felt (Union Carbide) hydrochloride (Aldrich) as the indicator. that was surface oxidized with boiling, concentrated nitric *Open-circuit oxidation of methanol by Fe(III) on a plati*acid for 15 min. This scheme may have advantages in the *nized platinum gauze electrode.* A two-compartment elecdesign of fuel cells. Particularly, it is not clear how to trochemical cell similar to one described previously (14)

design a catalytically active electrode that has a substantial **A number of alcohols, aldehydes, and carboxylic acids were** volume. However, with this scheme, it becomes difficult

*Materials*. Methanol, ethanol, and ethylene glycol were **INTRODUCTION** distilled before use. Acetic acid (glacial) was stirred with acetic anhydride and CrO<sub>3</sub> (2 g/100 ml) to remove water The catalytic oxidation of organic compounds using di-<br>oxygen as the ultimate oxidant is widely used in synthesis (Aldrich), glyoxyilic acid (50% solution, Aldrich), po-<br>and is a prevailing strategy for removal of contami



factor of 684, geometric area of  $6.25 \text{ cm}^2$ ) was prepared as trode was first carefully rinsed with a solution of  $30\%$  H<sub>2</sub>O<sub>2</sub> in concentrated H<sub>2</sub>SO<sub>4</sub> ( $\sim$ 10% H<sub>2</sub>O<sub>2</sub> solution in H<sub>2</sub>SO<sub>4</sub> by Ethylene glycol fuel cell. The assembly and operation volume) and then held at alternating potentials of  $+1.4$  of the cell was similar to that described for a methane and  $-0.1$  V in 1 M solutions of H<sub>2</sub>SO<sub>c</sub> in water at 25<sup>o</sup>C redox fuel cell (9). The oxidation of ethylen and  $-0.1$  V in 1 *M* solutions of H<sub>2</sub>SO<sub>4</sub> in water at 25<sup>o</sup>C until constant cyclic voltammograms (swept from 0 to 1.2 dispersed platinum black by Fe(III) and the reduction of V at 50 mV/s) were obtained. It was finally held at  $+1.4$  dioxygen by V(IV) were carried out in separate r V at 50 mV/s) were obtained. It was finally held at  $+1.4$ V for 20 s before being rinsed with water and transferred flasks. The anolyte and catholyte were circulated between<br>to the reaction vessel. The system was set up under an reformer flasks and cell compartments using perista to the reaction vessel. The system was set up under an experiment flasks and cell compartments using peristaltic atmosphere of argon with equal volumes of Fe(III) solution pumps. The solutions in the reformer flasks were f atmosphere of argon with equal volumes of Fe(III) solution pumps. The solutions in the reformer flasks were filtered<br>(0.6 M. Fe(III)/Pt<sub>argon</sub> = 978:1) and of 3 M H<sub>2</sub>SO<sub>4</sub> solution through glass frits before being admitt  $(0.6 M, Fe(III)/P_{\text{surface}} = 978:1)$  and of 3 *M* H<sub>2</sub>SO<sub>4</sub> solution through glass frits before being admitted to the fuel cell.<br>in the catalyst/electrode and reference electrode compart-<br>The reduction of Fe(III) to Fe(II) due to in the catalyst/electrode and reference electrode compartments, respectively. Control experiments showed that only ethylene glycol was carried out for 2 h before the solution slight variation in measured potentials of the catalyst/elec- was admitted to the anode compartment and the cell was trode occurred when the reference electrode was moved operated. Currents at various cell potentials were meabetween cell compartments. The system was allowed to sured under steady-state conditions by varying the load equilibrate in a  $60^{\circ}$ C bath before addition of methanol. across the cell. The cell compartments were 3 cm in diame-The potential of the working catalyst/electrode varied little ter and 5.5 cm deep and were separated by a Nafion memwith stir rate unless stirring was stopped, which would brane. A saturated calomel electrode was placed in each typically result in an increase of potential by  $\sim$ 50 mV. compartment as a reference electrode. Both the solutions

anol (see below), all reactions were performed as follows. chambers  $\Delta$ ll reactions were performed under argon using argon-cliftons. All reactions were performed under argon using argon-<br>nurged solutions After heating the platinum black [3 mol% The reformer flask for the anode consisted of a flask purged solutions. After heating the platinum black [3 mol%

allowing the reaction system to come to steady-state temperature (80 $^{\circ}$ C) and pressure (1 atm), the stopcock to a gas buret filled with mineral oil was opened. The organic compound to be oxidized was added as a neat liquid or as an aqueous solution via syringe. The amount added was selected so that the ratio of Fe(III) to reducing equivalents of fuel was approximately 2:1. Reaction generally occurred after a short  $(1-2 \text{ min})$  initiation period and was evidenced by gas production (an increase in the volume of gas in the buret). During the reaction, the volume of the system was allowed to expand by maintaining the pressure of the system at atmospheric by means of a sidearm reservoir on the gas buret. Reaction was carried out until FIG. 1. Schematic of the ethylene glycol/oxygen fuel cell. Details of gas production had slowed to virtually zero or after approxthe cell components and operation are provided in the Methods section. imately 3–4 h. Afterward, the reaction solution was filtered under argon atmosphere to remove the platinum black and titrated with a standard  $Ce^{4+}$  solution to determine the was used to study the oxidation. In this case, however, the<br>cell compartments were separated by a Nafion membrane,<br>a saturated calomel electrode was used as the reference<br>a rate that would interfere with the redox titratio electrode, and stirring was accomplished with a Teflon-<br>coated magnetic stirbar. The catalyst/electrode (roughness<br>imilar procedure except that 2 mol% of Pt based on the similar procedure except that 2 mol% of Pt based on the reducing equivalents of fuel was used, and the reaction previously described (15). The platinized catalyst/elec-<br>was carried out at a temperature of  $60^{\circ}$ C.

*Oxidation reactions.* Except for the oxidation of meth-<br>
let reformer flasks and the solutions in the fuel cell<br>
let only the reformed as follows chambers were stirred with Teflon-coated magnetic

based on the number of reducing equivalents of fuel (*x* in containing 75 ml of a 0.6 *M* Fe<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub> solution in 30% Eq. [1] and Table 1)], the flask was cooled to the reac-  $H_2SO_4$  solution, 1 ml of ethylene glycol [a four-fold excess tion temperature, and ferric sulfate solution (15 ml, relative to starting Fe(III)], and 79.8 mg of Pt black [0.9  $[Fe(III)] = 0.586$  *M*) was added. The stirring rate was mol% relative to starting Fe(III);  $3.3 \times 10^{-4}$  mol active established at 2400 rpm (1-in. Teflon-coated stirbar in a sites/g Pt as determined by  $H_2/O_2$  titrimetry (13) (6.4%) 100-ml, three-necked flask) using a stroboscope. After dispersion)]. The oxygen-reforming reaction is discussed

in detail elsewhere by Kummer and Oei (4–6). Cyclic volt- **TABLE 1** ammograms of the catholyte and anolyte were identical to<br>those of authentic solutions of  $V(V)$  and  $Fe(III)$ , respectively.

Both the anode and the cathode of the fuel cell consisted of rectangles of WDF graphite felt (Union Carbide,  $2 \times$  $3 \times 0.5$  cm). The felt was surface oxidized with boiling, concentrated nitric acid for 15 min. The density of the felt before this treatment was  $8.49 \times 10^{-2}$  g/cm<sup>3</sup>. The surface CH<sub>3</sub>OH 6 6.4 1 0.94 area and density of the felt after treatment with boiling nitric acid were not measured. The current collectors were Pt wires (0.5 mm in diameter and  $\sim$ 4 cm long). Since this redox fuel cell scheme permits the use of porous felt electrodes (i.e., the electrode reaction can take place in the volume of a porous electrode) and since the cost of the graphite felt is relatively unimportant, but the volume of the felt is important, current densities are reported per unit volume of carbon felt used (15).

Oxidation of methanol. Methanol is rapidly and quan-<br>titatively oxidized to  $CO_2$  by excess Fe(III) in the presence<br>dard  $Ce^{4+}$  solution and expressed as the ratio of  $[Fe^{2+}]$  at the end of the of a catalytic amount of platinum black (1 mol% Pt versus reaction to [*R*] at the beginning of the reaction. Error in this ratio is  $\pm 0.1$ . Fe(III);  $[Fe(HI)] = 0.7 M$ ;  $60^{\circ}$ C;  $30\%$  w:w  $H_2SO_4$  in water; <br>under argon). The rate of  $CO_2$  evolution was effectively temperature and pressure using a gas buret. Error in  $z_{\text{found}}$  is  $\pm 0.1$ . under argon). The rate of CO<sub>2</sub> evolution was effectively constant for the majority of reaction. The turnover frequency during this period was 0.39 turnover/atom  $Pt<sub>surface</sub>/$ s (one turnover = production of 1 CO<sub>2</sub> or of 6 Fe(III), Fe(III)  $\cong$  16:1) caused a rapid increase of catalyst/elec-<br>Pt<sub>surface</sub> is an active site on Pt black). Substantial evolution<br>of CO<sub>2</sub> was observed even when meth

mized (15) square of 52-mesh Pt gauze with a roughness<br>factor of 684 (15). The potential of the catalyst/electrode<br>was 1.15 V versus the standard hydrogen electrode (SHE;<br>all potentials in this paper are reported versus S sulted from the method we used to prepare the catalyst/ electrode surface (see the Methods section). Addition of  $R$  excess methanol (MeOH : Fe(III)  $\sim$  3.3 : 1) resulted in immediate reduction of the catalyst/electrode potential to 0.65 V and evolution of  $CO_2$  from the catalyst/electrode Table 1 summarizes the results of this reaction for various surface. The potential gradually decreased as Fe(III) was reductants (*R*). Our first interest was to establish the stoiconsumed, reaching a minimum (0.48 V versus SHE) at chiometries of these reactions. Reaction stoichiometry was the end of reaction, when evolution of  $CO<sub>2</sub>$  had ceased. quantified in two complementary ways: by measuring the Addition of more Fe(III) (remaining MeOH : added amount of  $CO<sub>2</sub>$  produced using a gas buret and by measur-





**RESULTS**<br> *a* Conditions 80°C, 3 *M* H<sub>2</sub>SO<sub>4</sub>.<br> *b* From Eq. [1].

of CO<sub>2</sub> was observed even when methanol was in excess We interpret this behavior as formation of the active cata-<br>of Fe(III) (e.g., MeOH: Fe(III) ~ 2:1; 12 reducing equiva-<br>lents per Fe(III), demonstrating that products

$$
R + yH_2O + xFe^{3+} \xrightarrow{Pt(0)} zCO_2 + xFe^{2+} + xH^+.
$$
 [1]



havior was observed and is illustrated in Fig. 3. ing the amount of ferric ion reduced to ferrous ion by redox titration. **DISCUSSION**

The first general class of compounds listed in Table 1 is  $C_2$  fuels in which each carbon atom is functionalized. These In this article, it has been shown that several alcohols, compounds were completely oxidized to  $CO_2$  at 80°C. The aldehydes and carboxylic acids were oxidize

Functionalized fuels with greater numbers of carbons gave results on oxidation that depended on the structure. Glycerin (three carbons) was oxidized completely under these reaction conditions. When six carbon substrates (glucose, sorbital) were employed as fuels, no gas evolved and only a small fraction of the expected ferrous ion was produced. Dehydrations of sugars in acidic media to form functionalized furans are documented in the literature (17). Given the results presented in this work, it is likely that such fuels are difficult to oxidize under these conditions and that cyclization/aromatization may be occurring.

The rates of oxidation of several fuels to carbon dioxide were monitored with a gas buret;  $CO<sub>2</sub>$  was identified by GC/MS. Figure 2 summarizes rates. Interestingly, a long<br>(20–30 min) initiation period was observed in the oxidation fuel cell. Anode and cathode voltages were measured using senarate of glycolic acid. This observation was made in triplicate reference electrodes in each compartment.

oxidations using different samples of substrate in each. Experimentally, there is ambiguity about the mechanisms of oxidation at a platinum surface (18). Empirically, multielectron oxidation is possible under these conditions from functionalized carbon compounds with no detectable slow steps.

*Operation of a fuel cell using ethylene glycol as the fuel.* Ethylene glycol was oxidized completely to carbon dioxide in a prototype fuel cell similar to one using methane (9) that was reported previously. Oxidation of ethylene glycol to  $CO<sub>2</sub>$  was carried out in a separate reformer flask, and the reduced iron solution was pumped into the anode chamber of the fuel cell to deliver the electron equivalents produced in the oxidation reaction. Since the redox couple **FIG. 2.** Rates of carbon dioxide production measured by gas buret. was the same as in the methane cell (9), the same electro-<br>Conditions are the same as those reported in Table 1. chemical behavior is expected if ethylene glycol does not oxidize much directly at the electrode. Indeed, similar be-

compounds were completely oxidized to  $CO_2$  at 80°C. The aldehydes, and carboxylic acids were oxidized under mild extent of this reaction after 3–4 h was, at a minimum, conditions (60–80°C 3 M H<sub>2</sub>SO<sub>4</sub> in water) by Fe(II extent of this reaction after 3–4 h was, at a minimum, conditions  $(60-80^{\circ}\text{C}, 3 \text{ } M \text{ H}_2\text{SO}_4)$  in water) by Fe(III) using 79% of completion as evidenced by the amount of CO<sub>2</sub> a catalytic amount of Pt black to gen 79% of completion as evidenced by the amount of  $CO<sub>2</sub>$  a catalytic amount of Pt black to generate Fe(II) and pro-<br>produced. By titration, it was found that at least enough tons Under these conditions complete oxidizat produced. By titration, it was found that at least enough tons. Under these conditions, complete oxidization of sev-<br>ferric ion was reduced to correspond to the expected eral reductants was effected On the basis of these r ferric ion was reduced to correspond to the expected eral reductants was effected. On the basis of these results,<br>a redox fuel cell that catalyzed the complete oxidation of mount of fuel oxidized (16).<br>
Ethanol and acetic acid both contain one unfunction-<br>
ethylene glycol by dioxygen at 80°C was assembled. A Ethanol and acetic acid both contain one unfunction-<br>alized (methyl) carbon. Under the experimental conditions<br>maximum power density of 9.9 mW/cm<sup>3</sup> of graphite felt alized (methyl) carbon. Under the experimental conditions maximum power density of 9.9 mW/cm<sup>3</sup> of graphite felt employed (80°C, 3–4 h reaction time), these reductants was obtained at a cell voltage of 197 mV. These resul employed (80°C, 3–4 h reaction time), these reductants was obtained at a cell voltage of 197 mV. These results were not completely oxidized to  $CO_2$ . No gas evolution indicate that the oxidation of the ethylene glycol in were not completely oxidized to  $CO<sub>2</sub>$ . No gas evolution indicate that the oxidation of the ethylene glycol in the was detected. The reduction of ferric ion suggests, howwas detected. The reduction of ferric ion suggests, how-<br>efformer section (Fig. 2) is not a limiting factor in the<br>ever, that partial oxidation of the ethanol occurred, proba-<br>electrical performance of the cell. The power ever, that partial oxidation of the ethanol occurred, proba-<br>bly to at least the oxidation level of acetic acid (expected this porous felt electrode is not straightforwardly compared bly to at least the oxidation level of acetic acid (expected this porous felt electrode is not straightforwardly compared  $x = 4$ ; see Eq [1]). We suspect that complete oxidation of with those in planar and gas diffusion e  $x = 4$ ; see Eq [1]). We suspect that complete oxidation of with those in planar and gas diffusion electrodes. However, these species can be effected, albeit under conditions of as the porous felt employed has substantial these species can be effected, albeit under conditions of as the porous felt employed has substantial surface area,<br>higher temperature and pressure (9).<br>it is clearly low compared to that in gas diffusion electrodes it is clearly low compared to that in gas diffusion electrodes



fuel cell. Anode and cathode voltages were measured using separate

(7, 8). However, by decoupling the catalytic oxidation of  $(7, 8)$ . However, by decoupling the catalytic oxidation of  $(7, 8)$ . Electrochem. 12, 41 (1982).<br>
the fuel from the electrochemical transfer of electrons to  $(985$ conventional, direct fuel cells. The electrode material is 8. Lindstrom, O., *Chemtech* **18**, 490 (1988).<br>constructed at substantially lower cost and with greater 9. Bergens, S. H., Gorman, C. B., Palmore, G. T. R., and Wh constructed at substantially lower cost and with greater 9. Bergens, S. H., Gorman, C. B., Pages than the conventional gas diffusion electrode that has G. M., Science 265, 1418 (1994). ease than the conventional gas diffusion electrode that has<br>been employed in a methanol fuel cell (19). As we have<br>shown previously, a high pressure and temperature re-<br> $\frac{C. M., Science 265, 1418 (1994)}{Corf. 13, 111 (1959)}$ .<br>The Sho former that catalyzes the complete oxidation of methane 12. Matsen, J. M., *Adv. Chem. Ser.* **64,** 277 (1967).<br>to carbon dioxide can be substituted for this reformer using 13. Carballo, L., Serrano, C., Wolf, E. E., and Ca to carbon dioxide can be substituted for this reformer using 13. Carballo, L., Serrano, C., Wolf, E. E., and Carberry, J. J., J. Catal.<br>the same cell scheme (9). Moreover, replacement and re-<br>covery of the catalyst is simp liquid reformers than for conventional fuel cell stacks (7). (A. J. Bard, Ed.), p. 1. Dekker, New York, 1976. Our results suggest that it is possible to develop redox fuel 16. In all cases but one, slightly more ferric ion was reduced than ex-<br>colle that oxidize a variety of fuels for both production of pected, resulting in a high cells that oxidize a variety of fuels for both production of<br>energy and for removal of contaminants from waste<br>streams; such work is in progress in our group.<br>treams; such work is in progress in our group.

a Postdoctoral Fellowship. **plete** oxidation of these substrates.

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- of Fe(III) was reduced, 4.45 mmol of which could be accounted for by complete oxidation of the glyoxal. If the rest of the iron oxidized the  $Pt(0)$ , it would correspond to complete oxidation of the  $Pt(0)$  to **ACKNOWLEDGMENTS**  $P_t(V)$ ,  $(4.92-4.45)/4 = 0.12$  mmol of  $P_t(0) \rightarrow P_t(V)$  but 24.3 mg  $Pt(0) \cdot (1 \text{ mmol}/195.09 \text{ mg}) = 0.12 \text{ mmol}$ . Furthermore, substantial This work was supported in part by the ONR and ARPA. SHB thanks amounts of platinum black remained after the reaction. Within this the National Science and Engineering Research Council of Canada for experimental uncertaint experimental uncertainty, this stochiometry does correspond to com-
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