Supplementary Material: Development of the Fray-Farthing-Chen Cambridge Process: Towards the Sustainable Production of Titanium and Its Alloys

DI HU^{1, 2, 4}, ALEKSEI DOLGANOV¹, MINGCHAN MA¹, BIYASH BHATTACHARYA¹, MATTHEW T. BISHOP^{1, 2}, and GEORGE Z. CHEN^{1, 2, 3, 4}

1.--Department of Chemical and Environmental Engineering, and Energy Engineering Research Group, Faculty of Science and Engineering, The University of Nottingham Ningbo China, Ningbo 315100, China.

2.--International Doctoral Innovation Centre, The University of Nottingham Ningbo China, Ningbo 315100, China.

3.--Department of Chemical and Environmental Engineering, and Advanced Materials Research Group, Faculty of Engineering, University of Nottingham, Nottingham NG7 2RD, UK.

4.--e-mails: di.hu@nottingham.edu.cn (Di Hu); george.chen@nottingham.ac.uk (George Z. Chen)

Table S1. Novel extraction technologies for titanium production [1-3]				
Name	Non-electrolytic process	Products		
The innovations in the Kroll Process itself [4]	in the Kroll (i) Fluidized bed chlorination; (ii) increased batch size; (iii) combined process technology; (vi) sponge handling and evaluation techniques; (v) development of energy-efficient cells			
Preform Reduction Process (PRP) [5]	Reduction of TiO ₂ by Ca vapour	Powder compact		
Mechanochemical Process [6]	Mechanical energy activated chemical reactions and structural changes. (i) Ambient temperature CaH ₂ reduction of TiO ₂ ; (ii) ambient temperature Mg reduction of TiCl ₄ .	Powder		
Armstrong Process [7]	Liquid Na reduction of TiCl ₄ vapor	Powder		
Name	Electrolytic process	Products		
Electronically Mediated Reaction (EMR) Process [8]	Electrolytic cell between TiO_2 and liquid Ca alloy reduces TiO_2	Highly porous Ti powder compact		
Materials and Electrochemical Research (MER) Process [9]	Dissolution of titanium cations based composite anode, transport through mixed halide electrolyte and deposition on cathode	Powder, flake or solid slab		
Ono-Suzuki (OS) Process [10]	Electrochemical-calciothermic reduction of TiO ₂	Powder / sponge		
Fray-Farthing-Chen-Cambridge Process (FFC-Cambridge Process) [11-12]	Electrolytic reduction of TiO ₂ electrode in molten salt	Powder block		
Solid Oxide Membrane (SOM) Process [13-14]*	Electrolytic reduction of dissolved Ti ions in a fluoride flux or solid TiO ₂ in a molten salt. Using oxygen ion conducting yttrium stabilised zirconia (YSZ) membrane to separate the anode from the molten electrolyte.	Deposited Ti / powder block		
Molten Oxide Electrolysis (MOE) Process [15-16]	Direct electrolysis of molten TiO ₂	Liquid Ti		

* For deoxidation of TiO₂, the cathodic reactions of the SOM process is identical to that of the FFC-Cambridge process, whereas its anodic reactions involve the removal of oxygen ions from the electrolyte via an oxide-ion-conducting solid electrolyte which separates the anodic reaction area from the molten electrolyte [13-14]. The carbon related side reactions are therefore confined within the tube of YSZ membrane, eliminating the carbon issues associated with the FFC-Cambridge process as described in the main text. However, because of the utilisation of an oxide-ion-conducting solid electrolyte, a high over-potential is therefore necessary, which results in a relatively higher cell voltage when compared with that of the FFC-Cambridge process (e.g. 3.8 V vs. 3.2 V for titanium production) [11, 14].



Fig. S1. Relative cost factors for conventional mill processing of a 2.54 cm thick Ti alloy plate (reprinted from [2] with permission).

The pathway of in situ perovskitisation [17]:

$5\text{TiO}_2(s) + \text{Ca}^{2+} + 2e^- = \text{Ti}_4\text{O}_7(s) + \text{Ca}\text{TiO}_3(s)$	RS1
$4\text{Ti}_4\text{O}_7(s) + \text{Ca}^{2+} + 2e^- = 5\text{Ti}_3\text{O}_5(s) + \text{Ca}\text{Ti}\text{O}_3(s)$	RS2
$3Ti_3O_5(s) + Ca^{2+} + 2e^- = 4Ti_2O_3(s) + CaTiO_3(s)$	RS3
$2Ti_2O_3(s) + Ca^{2+} + 2e^- = 3TiO(s) + CaTiO_3(s)$	RS4
$CaTiO_3(s) + TiO = CaTi_2O_4(s)$	RS5
$CaTi_2O_4(s) + 2e^- = 2TiO(s) + Ca^{2+} + 2O^{2-}$	RS6
$TiO(s) + 2(1-\delta)e^{-} = Ti_{[O]\delta} + (1-\delta)O^{2-}$	RS7

The electrochemical reduction of TiO_2 involves the consumption of CaO during the early stages, such as the incorporation of calcium ions into the oxide precursor (cf. **RS1** to **RS4**) meanwhile discharging oxygen ions at the anode, and production of CaO during the late stages, such as decomposition of calcium titanite (CaTi₂O₄) to titanium monoxide (TiO) and CaO (cf. **RS6**).

Pilling-Bedworth Ratio (PBR)

PBR =
$$V_o/V_m$$
E1 $V_o = M_o/n\rho_o$ E2 $V_m = M_m/\rho_m$ E3

Where the subscripts o and m represent the oxide and metal, respectively, V the molar volume, M the molar mass, ρ the density, and n the number of metal atoms in the oxide formula, e.g. n = 1 for TiO₂, n = 2 for aluminium oxide (Al₂O₃).



Fig. S2. An illustration shows the influence of PBR on electrochemical reduction of Cr₂O₃, TiO₂, and MgO, respectively (reproduced from [18] with permission).

As illustrated in **Fig. S2**, when the PBR is greater than unity, i.e. $V_0 > V_m$, a porous metal layer can be generated from the removal of oxygen. This porous layer is essential for the electrochemical reduction process as it could allow molten salt to access the underlying metal oxide. Thus, the reduction-generated oxygen ions (O²⁻) could be removed through the electrolyte in the pores of the metal layer, whist the solid state reduction of metal oxide to its metal could proceed (cf. chromium oxide (Cr₂O₃) to Cr as shown in **Fig. S2**, where PBR = 2). However, when the PBR is close to unity, the formed metal layer becomes less porous leading to a relatively low electrochemical reduction speed (cf. TiO₂)

to titanium in **Fig. S2**, where PBR = 1.33 based on TiO to titanium). When the PBR is much less than unity, deoxidation process will stall (cf. MgO to Mg in **Fig. S2**, where PBR = 0.81). It should be mentioned here, the final step in the process of electrochemical reduction of TiO₂ is TiO to titanium (as described in **RS7**), and thereby PBR for TiO to titanium is used to evaluate the kinetic difficulty level presented in the process.

Optimisation methods	Oxygen content in titanium (ppm by mass)	Energy consumption (kWh/kg Ti)	Current efficiency
Electrolyte composition ($CaCl_2 + 2 \mod CaO$), optimisation of polarisation regime (using a carbon pseudo-	3600	N/A	32%
porous TiO ₂ precursor (25 to 30% porosity), 16 hours of electrolysis [19].	2400	N/A	15%
Ex situ perovskitisation, porous $CaTiO_3$ (40 to 50% porosity), 10 hours of electrolysis [20].	2100	N/A	28%
Highly porous TiO ₂ precursor (68% porosity) achieved by using recyclable fugitive pore forming agent (NH ₄ HCO ₃), and optimised dual-stage for electrolysis (3.2 V, 3 hours + 2.6 V, 3 hours at 850 °C) [18].	1900	21.5	32.3%
Inert anode, 14 to 16 hours of electrolysis, and optimised electrolysis conditions [21].	N/A	17	40%

Table S2. A comparison of different optimisation methods developed for the FFC-Cambridge process for titanium extraction [18-21]

Chloride process (using purer ore or rutile as the feedstock):

$TiO_2 + C + 2Cl_2 = TiCl_4 + CO_2$	RS8a
$TiO_2 + 2C + 2Cl_2 = TiCl_4 + 2CO$	RS8b
$TiCl_4 + O_2 = TiO_2 + 2Cl_2$	RS9

Sulphate process (using ilmenite, i.e. FeTiO₃, as the feedstock):

$FeTiO_3 + 2H_2SO_4 = FeSO_4 + TiOSO_4 + 2H_2O$	RS10

 $TiOSO_4 + (n+1)H_2O = TiO_2 \cdot nH_2O + H_2SO_4$ RS11

RS12

$TiO_2 \cdot nH_2O = TiO_2 + nH_2O$



Fig. S3. Scanning electron microscopic images of (a) ground natural ilmenite feedstock, (b) its electrolysis product (electrolysis at 3.0 V, 900 °C, for 12 hours) (reprinted from [22] with permission), (c) synthetic rutile feedstock, and (d) backscattered image of consolidated (via spark plasma sintering) product from electro-reduction of synthetic rutile (with oxygen content ca. 3500 ppm) (reprinted from [23] with permission).



Fig. S4. (a) The current/oxygen-time plots for electrolysis of TiO₂ using a CaRuO₃ anode. Photos of (b) a pure CaRuO₃ anode after 150 hours electrolysis and washing, and (c, d) CaTi_xRu_{1-x}O₃ based inert anodes after electrolysis: (c) x = 0.72 and (d) x = 0.81 (reproduced from [24] with permission).

FFC-Cambridge process



Fig. S5. Ti-Ni alloy production by the FFC-Cambridge process and a conventional process

(reproduced from [25] with permission).

Commercial Progresses

Since 2001, MetalysisTM has been focusing on the commercialisation of the FFC-Cambridge process for titanium and other metals/alloys production. The titanium production capacity has been steadily increased from grams (GEN 1, R&D proof of concept, 2001 - 2003) to kilograms (GEN 2, Technical demonstrator, 2003 - 2009), then to tonnes (GEN 3, Commercial demonstrator, 2010 - 2014), and currently to 10s tonnes (GEN 4, Industrial scale production, 2015 - 2017) [26]. The GEN 5 facility will be the multiple modules of GEN 4, whose production capability can be easily adjusted from 100s to 1,000s tonnes according to different demands [26].

On the other hand, the FFC-Cambridge process has also been commercialised for production of silicon nanowire based negative electrode materials for Lithium-ion batteries by GLABATTM (a Chinese state-owned automotive battery research institute) [27].

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