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# 18

# **Understanding the Vanadium Redox Flow Batteries**

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# 1. Introduction

Vanadium redox flow batteries (VRB) are large stationary electricity storage systems with many potential applications in a deregulated and decentralized network. Flow batteries (FB) store chemical energy and generate electricity by a redox reaction between vanadium ions dissolved in the electrolytes. FB are essentially comprised of two key elements (Fig. 1): the cell stacks, where chemical energy is converted to electricity in a reversible process, and the tanks of electrolytes where energy is stored.

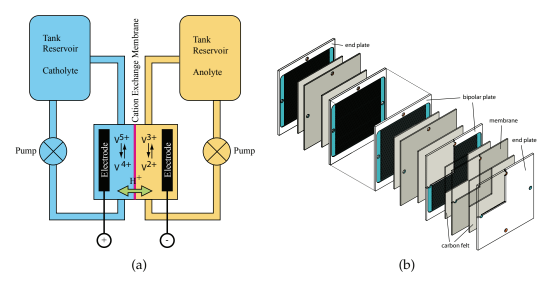


Fig. 1. (a) The schematics of the vanadium redox flow battery. (b) View of the different components composing a VRB stack. The surfaces in contact with the catholyte are coloured in blue and in orange for the anolyte.

The most significant feature of the FB is maybe the modularity of their power (kW) and energy (kWh) ratings which are independent of each other. In fact, the power is defined by the size and number of cells whereas the energetic capacity is set by the amount of electrolyte stored in the reservoirs. Hence, FB can be optimized for either energy and/or power delivery. Over the past 30 years, several redox couples have been investigated (Bartolozzi, 1989): zinc bromine, polysulfide bromide, cerium zinc, all vanadium, etc. Among them, VRB has the best chance to be widely adopted, thanks to its very competitive cost, its simplicity and because it contains no toxic materials.

In order to enhance the VRB performance, the system behaviour along with its interactions with the different subsystems, typically between the stack and its auxiliaries (i.e. electrolyte circulation and electrolyte state of charge), and the electrical system it is being connected to, have to be understood and appropriately modeled. Obviously, modeling a VRB is a strongly multidisciplinary task based on electrochemistry and fluid mechanics. New control strategies, based on the knowledge of the VRB operating principles provided by the model, are proposed to enhance the overall performance of the battery.

# 2. Electrochemistry of the vanadium redox batteries

Batteries are devices that store chemical energy and generate electricity by a reduction-oxidation (redox) reaction: i.e. a transformation of matter by electrons transfer. VRB differ from conventional batteries in two ways: 1) the reaction occurs between two electrolytes, rather than between an electrolyte and an electrode, therefore no electro-deposition or loss in electroactive substances takes place when the battery is repeatedly cycled. 2) The electrolytes are stored in external tanks and circulated through the stack (see Fig. 1). The electrochemical reactions occur at the VRB core: the cells. These cells are always composed of *a bipolar or end plate - carbon felt - membrane - carbon felt - bipolar or end plates*; they are then piled up to form a stack as illustrated in Fig. 1.

In the VRB, two simultaneous reactions occur on both sides of the membrane as illustrated in Fig. 2. During the discharge, electrons are removed from the anolyte and transferred through the external circuit to the catholyte. The flow of electrons is reversed during the charge, the reduction is now taking place in the anolyte and the oxidation in the catholyte.

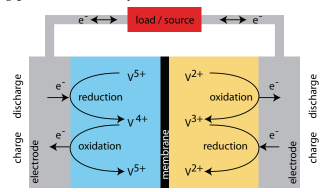


Fig. 2. VRB redox reaction during the charge and discharge

The VRB exploits the ability of vanadium to exist in 4 different oxidation states; the vanadium ions  $V^{4+}$  and  $V^{5+}$  are in fact vanadium oxide ions (respectively  $VO^{2+}$  and  $VO_2^+$ ). Thus, the VRB chemical equations become (Sum & Skyllas-Kazacos, 1985; Sum et al., 1985):

$$VO_{2}^{+} + 2H^{+} + e^{-} \rightleftharpoons VO^{2+} + H_{2}O$$

$$V^{2+} \rightleftharpoons V^{3+} + e^{-}$$

$$(1)$$

$$V^{2+} + VO_{2}^{+} + 2H^{+} \rightleftharpoons VO^{2+} + V^{3+} + H_{2}O$$

where the water ( $H_2O$ ) and protons ( $H^+$ ) are required in the cathodic reaction to maintain the charge balance and the stoichiometry.

#### 2.1 Equilibrium potential

The stack voltage  $U_{stack}$  depends on the equilibrium voltage  $U_{eq}$  and on the internal losses  $U_{loss}$ ; the equilibrium conditions are met when no current is flowing through the stack. In that case, there is no internal loss and  $U_{stack}$  equals  $U_{eq}$ ; otherwise, the internal losses modify  $U_{stack}$ . The internal losses<sup>1</sup>  $U_{loss}$  will be discussed in section 3.3. Hence  $U_{stack}$  is given by:

$$U_{stack}(t) = U_{eq}(t) - U_{loss}(t) [V]$$
<sup>(2)</sup>

The equilibrium voltage  $U_{eq}$  corresponds to the sum of the equilibrium potential E of the individual cells composing the stack. This potential is given by the Nernst equation and depends on the vanadium species concentrations and on the protons concentrations (Blanc, 2009):

$$E = E^{\Theta'} + \frac{RT}{F} \ln\left\{ \left( \frac{c_{VO_2^+} \cdot c_{H^+}^2}{c_{VO^{2+}}} \right) \left( \frac{c_{V^{2+}}}{c_{V^{3+}}} \right) \right\} \quad [V]$$
(3)

where *R* is the gas constant, *T* the temperature, *F* the Faraday constant,  $c_i$  the concentration of the species *i* and  $E^{e'}$  the formal potential. If we assume that the product/ratio of the activity coefficients is equal to 1, the formal potential  $E^{e'}$ , an experimental value often not available, can be replaced by the standard potential  $E^{e}$ .

#### 2.1.1 Standard potential from the thermodynamics

The standard potential  $E^{\diamond}$  is an ideal state where the battery is at standard conditions: vanadium species at a concentration of 1 M, all activity coefficients  $\gamma_i$  equal to one and a temperature of 25°C. The standard potential is an important parameter in the Nernst equation because it expresses the reaction potential at standard conditions; the second term in the Nernst equation is an expression of the deviation from these standard conditions. Together, they determine the equilibrium cell voltage under any conditions.

The standard potential  $E^{\oplus}$  can be found from thermodynamical principles, namely the Gibbs free enthalpy  $\Delta G$  and the conservation of energy, and empirical parameters found in electrochemical tables. We introduce here the standard Gibbs free enthalpy of reaction  $\Delta G^{\oplus}$  which represents the change of free energy that accompanies the formation of 1 M of a substance from its component elements at their standard states: 25°C, 100 kPa and 1 M (Van herle, 2002):

$$\Delta G^{\oplus} = \Delta H_r^{\oplus} - T \Delta S_r^{\oplus} \qquad [kJ/mol] \tag{4}$$

where the standard reaction enthalpy  $\Delta H_r^{\diamond}$  is the difference of molar formation enthalpies between the products  $\Delta H_{f,product}^{\diamond}$  and the reagents  $\Delta H_{f,reagent}^{\diamond}$ :

$$\Delta H_r^{\oplus} = \sum_{products} \Delta H_{f,product}^{\oplus} - \sum_{reagents} \Delta H_{f,reagent}^{\oplus} \qquad [kJ/mol]$$
(5)

and the standard reaction entropy  $\Delta S_r^{\diamond}$  is the difference of molar formation entropies between the products  $S_{f,vroduct}^{\diamond}$  and the reagents  $S_{f,reagent}^{\diamond}$ :

$$\Delta S_r^{\oplus} = \sum_{products} S_{f,product}^{\oplus} - \sum_{reagents} S_{f,reagent}^{\oplus} \qquad [J/mol \cdot K]$$
(6)

<sup>1</sup>Note that the sign of  $U_{loss}$  depends on the operating mode (charge or discharge).

Then, when we introduce the thermodynamical data from Tab. 1 into (5), the standard reaction enthalpy  $\Delta H_r^{\ominus}$  of the VRB reaction (1) becomes:

$$\Delta H_r^{\oplus} = \Delta H_{f,VO^{2+}}^{\oplus} + \Delta H_{f,V^{3+}}^{\oplus} + \Delta H_{f,H_2O}^{\oplus} - \Delta H_{f,V^{2+}}^{\oplus} - \Delta H_{f,VO_2^{\pm}}^{\oplus} - 2\Delta H_{f,H^{+}}^{\oplus}$$

$$= -155.6 \, kJ/mol$$
(7)

and similarly, the standard reaction entropy  $\Delta S_r^{\diamond}$  is obtained when these thermodynamical data are introduced into (6):

$$\Delta S_r^{\oplus} = S_{f,VO^{2+}}^{\oplus} + S_{f,V^{3+}}^{\oplus} + S_{f,H_2O}^{\oplus}$$
$$- S_{f,V^{2+}}^{\oplus} - S_{f,VO_2^{\pm}}^{\oplus} - 2S_{f,H^{\pm}}^{\oplus}$$
$$= -121.7 \ J/mol \cdot K$$
(8)

Formula	State	$\Delta H_{f}^{\ominus}\left[kJ/mol ight]$	$\Delta G_{f}^{\oplus}\left[kJ/mol\right]$	$S_f^{\oplus}\left[J/mol\cdot K\right]$
$V^{2+}$	aq	(-226)	-218	(-130)
$V^{3+}$	aq	(-259)	-251.3	(-230)
$VO^{2+}$	aq	-486.6	-446.4	-133.9
$VO_2^+$	aq	-649.8	-587.0	-42.3
$H_2O$	aq	-285.8	-237.2	69.9
$H^+$	aq	0	0	0

Table 1. Thermodynamical data for some vanadium compounds at 298.15 K. Values in parentheses are estimated (Van herle, 2002; Bard et al., 1985).

The conservation of energy relates the change in free energy resulting from the transfer of n moles of electrons to the difference of potential E:

$$\Delta G = -nFE \qquad [J/mol] \tag{9}$$

Therefore, we obtain the standard potential  $E^{\oplus}$  when we introduce  $\Delta G^{\oplus}$  (4) with the values of the standard reaction enthalpy (7) and entropy (8) into the reformulated (9):

$$E^{\oplus} = -\frac{\Delta G^{\oplus}}{nF} = -\frac{\Delta H_r^{\oplus} - T\Delta S_r^{\oplus}}{nF} \qquad [V]$$
(10)

So, we have determined from the thermodynamical principles that the standard potential  $E^{\oplus}$  is 1.23 V at 25°C.

The characteristic curve of the equilibrium potential *E* is illustrated in Fig. 3 for a single cell as a function of the state of charge *SoC*. We can also observe the relation between *E*, *SoC* and the protons and vanadium concentrations.

	Salt	Charge	Discharge	Electrolyte
$V^{2+}$	$VSO_4$	$\uparrow$	$\downarrow$	Anolyte
$V^{3+}$	$0.5 V_2(SO_4)_3$	$\downarrow$	$\uparrow$	Anolyte
$V^{4+}$ or $VO^{2+}$	$VOSO_4$	$\downarrow$	$\uparrow$	Catholyte
$V^{5+}$ or $VO_2^+$	$0.5 (VO_2)_2 SO_4$	$\uparrow$	$\downarrow$	Catholyte

Table 2. The different vanadium ions with their corresponding salt, their concentration variation during the charge and discharge of the VRB, and the electrolyte where they are dissolved.

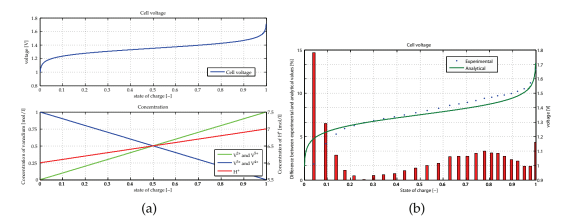


Fig. 3. (a) Top: Cell voltage versus the state of charge at  $25^{\circ}$ C. Bottom: Protons  $H^+$  and vanadium concentrations. (b) Comparison between the Nernst equation (3) and the experimental data published in (Heintz & Illenberger, 1998). The red bars represent the difference between the analytical and experimental data.

# 3. Electrochemical model

The main electrochemical relations governing the equilibrium voltage where introduced in the previous section. In order to have an electrochemical model of the VRB, it is now necessary to describe how the vanadium concentrations vary during the battery operation.

# 3.1 Concentration of vanadium ions

We see clearly from (1) that during the redox reactions, the vanadium ions are transformed and that some protons  $H^+$  are either produced or consumed. Therefore, the ion concentrations must change in the electrolyte to reflect these transformations which depend on how the battery is operated.

For example, when the battery is charged,  $V^{2+}$  and  $VO_2^+$  are produced and their concentrations increase; and  $V^{3+}$  and  $VO^{2+}$  are consumed and thus their concentrations diminish. This process is reversed when the battery is discharged. Tab. 2 summarizes the direction of the change for each species.

#### 3.1.1 Electron exchange rate

Obviously, the concentration changes are proportional to the reaction rate; and from (1) we also know that an electron is involved each time a redox reaction occurs. Therefore, the concentration changes are also proportional to the electrical current. Thus, the pace of the concentration variation is set by the electrical current flowing through the cell:

$$Q_c = n_{e^-}e = \int i(t)dt \qquad [C] \tag{11}$$

where  $Q_c$  is the charge, *i* the current, *t* the time,  $n_{e^-}$  the number of electrons and *e* the elementary charge. Therefore, the number of electrons  $n_{e^-}$  involved for a given current<sup>2</sup> is:

$$n_{e^-} = \frac{1}{eN_A} \int i(t)dt \qquad [mol] \tag{12}$$

where  $N_A$  is the Avogadro number. Then (12) leads to the definition of a molar flowrate of electrons  $\dot{N}_{e^-}$ :

$$\dot{N}_{e^-}(t) = \frac{1}{eN_A}i(t) \qquad [mol/s]$$
(13)

Physically, an electron is released by the oxidation of a vanadium ion, travels through the electrodes and is captured by the reduction of another vanadium ion in the opposite half-cell. In the case of a stack composed of  $N_{cell}$  cells, the electrons travel through the bipolar electrode to the adjacent cell (Fig. 4). Thus, for one electron flowing through the external electrical circuit,  $N_{cell}$  redox reactions have occurred. Therefore, the total molar flowrate of electrons  $\dot{N}_{e_{int}}$  for a stack is obtained by multiplying (13) by the number of cells:

$$\dot{N}_{e_{tot}^-}(t) = \frac{N_{cell}}{eN_A}i(t) = \frac{N_{cell}}{F}i(t) \qquad [mol/s]$$
(14)

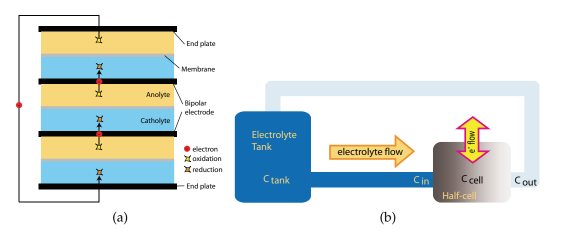


Fig. 4. (a) Illustration of the redox reactions required to produce a one electron flow in a 3 elements stack during the discharge. When the battery is charged, the flow and the reactions are inverted. (b) Illustration of the hydraulic circuit (half cell) where the concentrations are shown.

<sup>&</sup>lt;sup>2</sup>By convention, the current is positive during the VRB discharge in order to have a positive power delivered by the battery.

### 3.1.2 Input, output and average concentrations of vanadium ions

We know now that the vanadium concentrations change within the cells when the battery is operating. Therefore, the concentrations are not uniformly distributed through the electrolyte circuit (Fig. 4). Indeed, four concentrations are located in the VRB: the tank concentration  $c_{tank}$ , the concentration at the cell input  $c_{in}$ , the concentration inside the cell  $c_{cell}$  and the concentration at the cell output  $c_{out}$ .

Usually, the size of the reservoir is large compared to the electrolyte flowrate; thus the change in concentrations due to the flow of *used* electrolyte is so small that the tank concentrations are considered homogeneous. And therefore, the input concentrations  $c_{in}$  correspond exactly to  $c_{tank}$ .

The tank concentration  $c_{tank}$  reflects the past history of the battery; indeed the change in  $c_{tank}$  is proportional to the quantity of vanadium that has been transformed in the stack: this value corresponds to the quantity of electrons involves in the reaction. Therefore,  $c_{tank}$  is defined by the initial ion concentrations  $c_{tank_i}^{initial}$ , the size of the reservoir  $V_{tank}$  and the total molar flowrate of electrons  $\dot{N}_{e_{tan}}$ :

$$c_{in_{i}}(t) = c_{tank_{i}}(t) = c_{tank_{i}}^{initial} + \frac{1}{V_{tank}} \int b \dot{N}_{e_{tot}}(t) dt$$

$$= c_{tank_{i}}^{initial} + \frac{1}{V_{tank}} \int \frac{b}{F} i(t) dt \qquad [mol/l]$$
(15)

where *b* is a sign factor that reflects the direction of the reaction in accordance with Tab. 2:

$$b = \begin{cases} -1 \text{ for } V^{2+} \text{ and } V^{5+} \text{ ions} \\ 1 \text{ for } V^{3+} \text{ and } V^{4+} \text{ ions} \end{cases} [-]$$
(16)

The description of the output concentration  $c_{out}$  is difficult because it depends on the electrolyte flowrate Q, the length of the electrolyte circuit and on the current i that the electrolyte encounters during the cell crossing. Since the distribution of the vanadium ions inside the cell is unknown, we consider that the model has no memory and reacts instantly to a change in the operating conditions. In that case,  $c_{out}$  is related to the electrons molar flowrate  $\dot{N}_{e_{int}}$ , the electrolyte flowrate Q and on the input concentration  $c_{in}$ :

$$c_{out_i}(t) = c_{in_i}(t) + b \frac{N_{e_{tot}}(t)}{Q(t)} = c_{in_i}(t) + \frac{bN_{cell}}{F} \frac{i(t)}{Q(t)} \qquad [mol/l]$$
(17)

where: 
$$c_i$$
 = concentration of the different vanadium ions  $[mol/l]$   
 $Q(t)$  = flowrate of the electrolyte  $[l/s]$ 

For a quasi steady state, where the current and the flowrate are almost constant, the model predicts accurately the output concentrations. Unfortunately, it is not able to predict the transient behaviour when the system encounters extreme conditions such as the combination of a low flowrate, few active species and sudden current change. But when these conditions are avoided, (17) offers a very good insight of the battery behaviour.

We still have to establish the most important concentration: the concentration inside the cell  $c_{cell}$  that is necessary to solve the Nernst equation (3). Because the ion concentrations are not uniformly distributed inside the cell, we will make an approximation to determine  $c_{cell}$  from the mean value of  $c_{in}$  and  $c_{out}$ :

$$c_{cell_i}(t) = \frac{c_{in_i}(t) + c_{out_i}(t)}{2}$$
 [mol/l] (18)

#### 3.2 Concentration of protons

Unfortunately, (1) does not reflect exactly the phenomena happening in the cells. Indeed, the VRB electrolytes contain not only vanadium ions at different oxidation states, but also protons  $H^+$  and sulphate ions  $SO_4^{2-}$  that are only partially represented in the chemical equations; these ions are called spectator ions and do not take an active part in the reaction. But these spectator ions are important to respect the law of conservation of mass and the charge balance in both electrolytes (Blanc, 2009). The complete ionic equation, illustrated in Fig. 5, is useful to understand how the protons concentration  $c_{H^+}$  changes and why the protons cross the membrane to balance the charge.

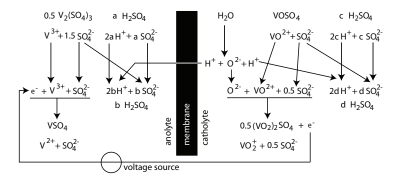


Fig. 5. Illustration of the full ionic equations of the VRB during the charge.

Hence, the protons concentration in the catholyte depends on the electrolyte composition and varies with the state of charge:

$$c_{H^+} = c_{H^+,discharged} + c_{VO^{2+}} [M]$$
<sup>(19)</sup>

where  $c_{H^+,discharged}$  is the protons concentration when the electrolyte is completely discharged.

#### 3.3 Internal losses

When a net current is flowing through the stack, the equilibrium conditions are not met anymore and the stack voltage  $U_{stack}$  is now given by the difference between the equilibrium potential  $U_{eq}$  and the internal losses  $U_{loss}$ . These losses are often called overpotentials and represent the energy needed to force the redox reaction to proceed at the required rate; a list of the variables affecting this rate is given in Fig. 6.

$$U_{loss}(t) = \eta_{act}(t) - \eta_{conc}(t) - \eta_{ohm}(t) - \eta_{ion}(t) [V]$$
<sup>(20)</sup>

The activation  $\eta_{act}$  and the concentration  $\eta_{conc}$  overpotentials are electrode phenomena and are respectively associated with the energy required to initiate a charge transfer and caused by concentration differences between the bulk solution and the electrode surface; in addition, the ohmic  $\eta_{ohm}$  and ionic  $\eta_{ionic}$  losses also alter the stack voltage. The ohmic losses  $\eta_{ohm}$  occur in the electrodes, the bipolar plates and the collector plates and the ionic losses  $\eta_{ionic}$  occur in the electrolytes and the membranes. But these overpotentials are seldom found in the literature and often applicable only to peculiar conditions. Therefore, an equivalent resistance is introduced instead:

$$U_{loss}(t) = R_{eq,charge/discharge}i(t) \ [V]$$
(21)

where  $R_{eq,charge}$  is the equivalent charge resistance and  $R_{eq,discharge}$  corresponds to the discharge resistance; these values are found experimentally (Skyllas-Kazacos & Menictas, 1997) and depends on the electrolyte, electrode materials and stack construction.

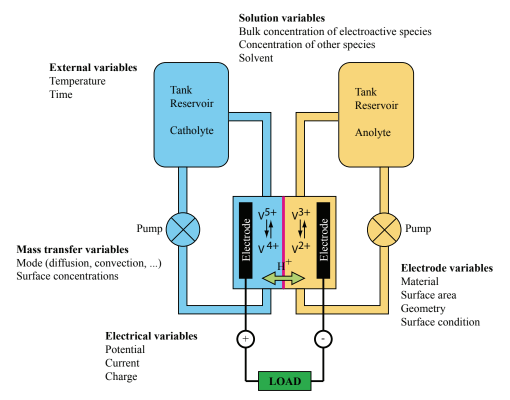


Fig. 6. Schematic representation of VRB with a list of variables affecting the rate of the redox reaction (Bard & Faulkner, 2001). Note that only one cell is represented on this figure.

#### 3.4 State of charge

The state of charge *SoC* indicates how much energy is stored in the battery; it varies from 0 (discharged state) to 1 (charged) and is defined by the following relation:

$$SoC = \left(\frac{c_{V^{2+}}}{c_{V^{2+}} + c_{V^{3+}}}\right) = \left(\frac{c_{VO_2^+}}{c_{VO^{2+}} + c_{VO_2^+}}\right) [-]$$
(22)

#### 3.5 Electrochemical model

From the principles explained in the previous section, it is now possible to introduced the electrochemical model that describes the behaviour of the stack, mainly how the stack voltage  $U_{stack}$  depends on the operating conditions: the current *I*, the vanadium concentrations in the electroactive cells  $c_{cell}$ , the protons concentration  $c_{H^+}$ , the electrolyte flowrate *Q* and the temperature *T*; furthermore, it also describes how the electrolyte compositions change as the battery is operating. The schematic representation of this model is shown in Fig. 7.

#### 3.6 Efficiencies

Efficiencies are parameters used to assess the performance of storage system. Basically, the definition of efficiency is simple, the energy efficiency  $\eta_{energy}$  is the ratio of the energy furnished by the battery during the discharge to the energy supplied during the charge:

$$\eta_{energy} = \frac{\int P_{VRB,discharge}(t)dt}{\int \left| P_{VRB,charge} \right|(t)dt} \qquad [-]$$
(23)

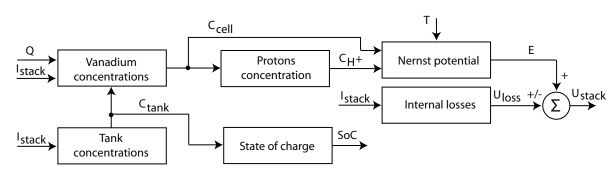


Fig. 7. Schematic representation of the electrochemical model

Name	Value	Name	Value
number of cells N <sub>cells</sub>	19	electrolyte vanadium concentration	2 M
R <sub>charge</sub>	$0.037 \ \Omega$	tank size $V_{tk}$	831
R <sub>discharge</sub>	0.039 Ω	initial concentration of vanadium species	1 M
electrolyte flowrate Q	21/s	_	

Table 3. The characteristics of the VRB stack.

But difficulties quickly arise when different technologies or products are compared because the operating mode has a significant impact on the performance: a quick charge produces more losses than a gentle one. The coulombic efficiency  $\eta_{coulombic}$  is a measure of the ratio of the charge withdrawn from the system  $Q_{discharge}$  during the discharge to the charge  $Q_{charge}$ supplied during the charge:

$$\eta_{coulombic} = \frac{Q_{discharge}}{Q_{charge}} = \frac{\int i_{discharge}(t)dt}{\int \left|i_{charge}(t)\right|dt} \qquad [-]$$
(24)

The voltage efficiency  $\eta_{voltage}$  is defined for a charge and discharge cycle at constant current. It is a measure of the ohmic and polarisation losses during the cycling. The voltage efficiency is the ratio of the integral of the stack voltage  $U_{stack,discharge}$  during the discharge to that of the voltage  $U_{stack,charge}$  during the charge:

$$\eta_{voltage} = \frac{\int U_{stack,discharge}(t)dt}{\int U_{stack,charge}(t)dt} \neq \frac{\eta_{energy}}{\eta_{coulombic}} \quad [-]$$
(25)

Note that when the mechanical losses  $P_{mech}$  are taken into account,  $\eta_{voltage}$  is not equal to the ratio of  $\eta_{energy}$  to  $\eta_{coulombic}$ .

## 3.7 Charge and discharge cycles at constant current

The electrochemical model of the vanadium redox battery is compared in this section to experimental data. To determine the performance, a VRB composed of a 19 elements stack and two tanks filled with 83 l of electrolytes will be used. The total vanadium concentration in each electrolyte is 2 M. The characteristics of the stack are summarized in Tab. 3 and correspond to an experimental stack built by M. Skyllas-Kazacos and co-workers (Skyllas-Kazacos & Menictas, 1997). The electrochemical model is used to assess the stack efficiencies during a series of charge and discharge cycles at constant currents.

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Current [A]	η <sub>energy</sub> [%]	η <sub>voltage</sub> [%]	η <sub>coulombic</sub> [%]	Current [A]	η <sub>energy</sub> [%]	η <sub>voltage</sub> [%]	η <sub>coulombic</sub> [%]
SI	SIMULATION RESULTS			EXP	'ERIMEN'	TAL DAT	A
10	97.0	97.0	100				
20	94.1	94.1	100				
40	88.6	88.6	100	60	78.3	82.8	94.6
60	83.3	83.3	100	100 (cycle 1)	68.0	72.3	94.0
80	78.4	78.4	100	100 (cycle 2)	70.8	73.0	96.1
100	73.7	73.7	100	100 (cycle 3)	73.1	74.0	98.7

Table 4. Efficiencies at various currents. The cycle starts at 2.5% SoC, the battery is charged until a 97.5% SoC and then discharged until a 2.5% SoC. Experimental data are from (Skyllas-Kazacos & Menictas, 1997).

At the beginning of the cycle, the battery state of charge *SoC* is 2.5% (discharged); the battery is charged at constant current until a *SoC* of 97.5% and then discharged until it reached its initial *SoC*. The resulting stack voltages  $U_{stack}$  and power  $P_{stack}$  are illustrated in Fig. 8 and the efficiencies are summarized in Tab. 4 along with experimental data. We observe quickly that the efficiencies decrease as the current increases.

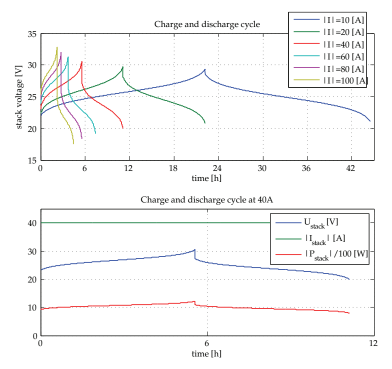


Fig. 8. Stack voltages during charge/discharge cycles at diverse currents. Below: stack voltage, current and power at 40A.

The voltage efficiencies  $\eta_{voltage}$  are accurately determined by the model; the difference with the experimental data always stays below 2%. The losses in coulombic efficiency  $\eta_{coulombic}$  can be caused by side reactions or cross mixing of electrolyte through the membrane which are not taken into account in the model; note that  $\eta_{coulombic}$  has improved as the battery becomes conditioned. When  $\eta_{coulombic}$  is close to 100%, as it is the case for the last cycle,

Solution of vanadium	Density [g/cm <sup>3</sup> ]	Viscosity [cP]	Vanadium / sulphuric acid concentration [M]
$V^{2+}$	1.2-1.3	1.7-2.4	1-2 / 2
$V^{3+}$	1.2-1.5	1.7-9.6	0.5-3 / 2
$V^{4+}$		(3.6-33.7)	0.25-3 / 3
$V^{4+}$	1.2-1.5		1-2 / 1-9
$V^{5+}_{V^{5+}}$	1.2-1.5		1-2 / 1-9
$V^{5+}$		3.2-22.3	0.5-3 / 4-7

Table 5. Density and viscosity of vanadium species solutions (Mousa, 2003; Wen et al., 2006; Oriji et al., 2004; Kausar, 2002). The numbers in brackets are estimations made from the kinematic viscosity.

the experimental and simulated energy efficiencies  $\eta_{energy}$  are almost the same, the difference being less than 1%. In the worst case, cycle 1, the difference is around 8.3%.

# 4. Electrolyte properties

The electrolyte properties are important parameters in the mechanical model; the density indicates its inertia, or resistance to an accelerating force, and the viscosity describes its *fluidity*, it may be thought of as internal friction between the molecules. They are both related to the attraction forces between the particles; thus they depend on the electrolyte composition.

The VRB electrolytes are composed of vanadium ions dissolved in sulphuric acid; we have seen previously that their composition changes as the battery is operating (see Fig. 3). Therefore, the electrolyte properties must change accordingly to the composition; but for simplicity reasons, these properties are maintained constant in this work. Tab. 5 gives the density and the viscosity for some vanadium solutions.

# 5. Fluid mechanics applied to the vanadium redox flow batteries

We introduce in this section the mechanical model that determines the power  $P_{pump}$  required to flow the electrolytes from the tanks through the stack and back in the tanks (see Fig. 1). This model is composed of an analytical part that models the pipes, bends, valves and tanks and a numerical part that describes the more complex stack hydraulic circuit.

# 5.1 Hydraulic circuit model (without the stack)

The analytical hydraulic model describes the pressure drop  $\Delta p_{pipe}$  in the pipes, the valve and the tank; it is based on the extended Bernoulli's equation that relates  $\Delta p_{pipe}$  to the fluid velocity  $V_s$ , the height z, the head loss  $h_f$  due to the friction and the minor losses  $h_m$ :

$$\Delta p_{pipe} = -\gamma \left( \frac{\Delta V_s^2}{2g} + \Delta z + h_f + h_m \right) \qquad [Pa]$$
<sup>(26)</sup>

where  $\gamma$  is the specific weight and g the gravitational acceleration. The head losses are obtained by dividing the hydraulic circuit into small

The head losses are obtained by dividing the hydraulic circuit into smaller sections where  $h_{f,i}$  or  $h_{m,i}$  are easily determined with the Darcy-Weisbach equation (Munson et al., 1998):

$$h_{f,i} = f_i \frac{L_i}{D_i} \frac{V_{s,i}^2}{2g}, \quad h_{m,i} = k_{L,i} \frac{V_{s,i}^2}{2g} \qquad [m]$$
 (27)

geometry	Loss coefficient $k_{L,i}$
from a reservoir into a pipe	0.04 - 0.9
from a pipe into a reservoir	1
bends and elbows	0.2 - 1.5
valves	0.15 - 10

Table 6. Loss coefficients (Munson et al., 1998; Candel, 2001).

where  $f_i$  is the friction factor,  $k_{L,i}$  the loss coefficient given in Tab. 6,  $L_i$  and  $D_i$  are the length and diameter of the conduit.

When the flow is laminar, the friction factor  $f_i$  is derived from the Poiseuille law (28) and for a turbulent flow, it is obtained from the Colebrook equation (29) (Candel, 2001):

$$f_i = \frac{64}{Re_i} \qquad [-] \tag{28}$$

$$\frac{1}{\sqrt{f_i}} = -2\log\left(\frac{\epsilon_i}{3.7D_i} + \frac{2.51}{Re_i\sqrt{f_i}}\right) \qquad [-]$$
<sup>(29)</sup>

where  $\epsilon_i$  is the equivalent roughness of the pipe and  $Re_i$  is the Reynolds number:

$$Re = \frac{\rho V_s D}{\mu} = \frac{V_s D}{\nu} \qquad [-] \tag{30}$$

where  $\rho$  is the density,  $\mu$  the dynamic viscosity and  $\nu$  the kinematic viscosity.

## 5.2 Stack hydraulic model

The stack geometry is too complex to be analytically described (Fig. 9), therefore the stack hydraulic model can only be numerically obtained with a finite element method (FEM).

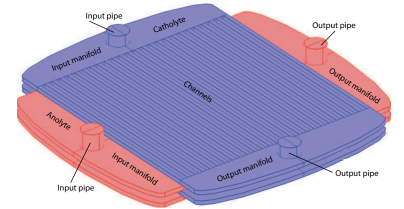


Fig. 9. Hydraulic circuit of a 2 cells stack. Note that the frame is not represented and that the colored segments represented the electrolytes (liquid).

It was assumed that the flow stays laminar in the stack; although the flow might be turbulent in the manifold at high velocity. In this example, the flow stays laminar in the distribution channels where the major part of the pressure drop  $\Delta p_{stack}$  occurs; therefore, the pressure drop in the stack  $\Delta p_{stack}$  is proportional to the flowrate:

$$\Delta p_{stack} = Q\tilde{R} \qquad [Pa] \tag{31}$$

where  $\vec{R}$  is the hydraulic resistance obtained from FEM simulations.

#### 5.3 Mechanical model

Finally, the sum of the pressure drop in the pipes  $\Delta p_{pipe}$  and the pressure drop in the stack  $\Delta p_{stack}$  determines the hydraulic circuit pressure drop  $\Delta p_{system}$ :

$$\Delta p_{system} = \Delta p_{vive} + \Delta p_{stack} \qquad [Pa] \tag{32}$$

The pump power  $P_{pump}$ , a determinant variable that influences the battery performance is related the head rise  $h_p$  supplied by the pump, to the fluid density  $\gamma$  and to the flowrate Q; we can also relate it to the pressure drop  $\Delta p$  (Wilkes, 2005):

$$P_{pump} = \gamma h_p Q = \Delta p \ Q \qquad [W] \tag{33}$$

The efficiency of the pump  $\eta_{pump}$  is affected by the hydraulic losses in the pump, the mechanical losses in the bearings and seals and the volumetric losses due to leakages inside the pump. Although  $\eta_{pump}$  is not constant in reality, it is assume in this work. Therefore, the effective power required by the pump  $P_{mech}$  is given by:

$$P_{mech} = \frac{P_{pump}}{\eta_{pump}} \qquad [W] \tag{34}$$

Thus, the relations introduced in this section can be combined to form the mechanical model of the VRB as illustrated in Fig. 10. Remember that the VRB needs two pumps to operate.

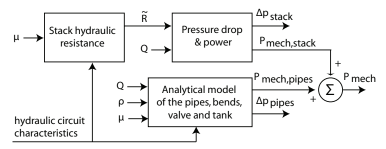


Fig. 10. Flowchart of the VRB mechanical model.

# 6. Multiphysics model and energetic considerations

The combination of the electrochemical model and the mechanical model leads to the multiphysics VRB system model. The functions that determine the vanadium concentrations in the tank  $c_{tank}$  and the state of charge *SoC* have been separated from the electrochemical model in order to be incorporated into a new model named *reservoir and electrolyte model*. A system control has also been added to supervise the battery operation; this system controls the flowrate Q and the stack current  $I_{stack}$ . This multiphysics system model, illustrated in Fig. 11, is a powerful means to understand the behaviour of the VRB, identify and quantify the sources of losses in this storage system; thus this multiphysics model is a good means to enhance the overall VRB efficiency.

#### 6.1 Power flow

In order to optimize the performance of the VRB, it is important to understand the power flows within the VRB storage system. The power converters represented in Fig. 12 are necessary to adapt the stack voltage  $U_{stack}$  to the power source  $U_{grid}$  or to the load voltage  $U_{load}$  and to supply the mechanical power required to operate the pumps. Since power

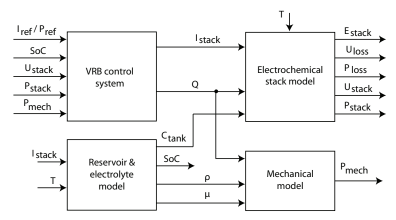


Fig. 11. Structured diagram of the multiphysics VRB system model.

converters are very efficient, with efficiencies around 98 to 99% (Wensong & Lai, 2008; Burger & Kranzer, 2009), they are considered, for simplicity, lossless in this work. Therefore, they are two sources of losses: the internal losses that are already included in the stack voltage  $U_{stack}$  (2), and the mechanical losses  $P_{mech}$ . Hence,  $P_{mech}$  is provided from the external power source during the charge and from the stack during the discharge. By convention, the battery power  $P_{VRB}$  and the stack power  $P_{stack}$  are positive during the discharge and negative during the charge;  $P_{mech}$  is always positive. Thus,  $P_{VRB}$  is given by:

$$P_{VRB} = P_{stack} - P_{mech} \qquad [W] \tag{35}$$

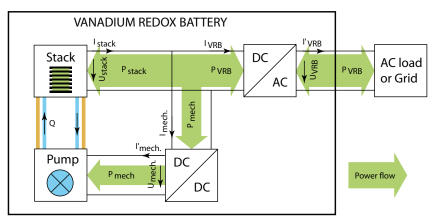


Fig. 12. Power flow in the VRB storage system. In this example, the power converters only adapt the currents and voltages, and are lossless.

In the rest of this section, we will discuss the battery performance under different operating strategy with a strong focus on the battery power  $P_{VRB}$ , the stack power  $P_{stack}$  ant the mechanical power  $P_{mech}$ . Intuitively, we feel that there should be an optimal control strategy that maximizes the battery performance. In these circumstances, the power delivered to the battery at any operating point is minimized during the charge and the power supplied by the battery is maximized during the discharge.

# 7. Operation at maximal and minimal flowrates

First, we will discuss the battery operation at maximal and minimal flowrates. We must keep in mind that an efficient control strategy must maximize the power exchanged with the battery

Name	Value
number of cells N <sub>cells</sub>	19
R <sub>charge</sub>	$0.037~\Omega$
R <sub>discharge</sub>	0.039 Ω
flow resistance $\widetilde{R}$	14186843 Pa/m <sup>3</sup>
electrolyte vanadium concentration	2 M
tank size $V_{tank}$	83 1
initial concentration of vanadium species	1 M

Table 7. the parameters of the simulation.

while minimizing the losses; there is no point to have a battery that consumes more power than necessary. To illustrate this discussion, we will use a 2.5 kW, 6 kWh VRB in the rest of this chapter; its characteristics are summarized in Tab. 7.

#### 7.1 Maximal flowrate

The simplest control strategy operates the battery at a constant flowrate set to provide enough electroactive species to sustain the chemical reaction under any operating conditions. Therefore, this flowrate  $Q_{max}$  is determined by the worst operating conditions: low state of charge *SoC* during the discharge and high *SoC* during the charge at high current in both cases. For the battery described in Tab. 7,  $Q_{max}$  is around 1.97 l/s: in that case, the mechanical power  $P_{mech}$  is 1720 W. In order to assess the performance, an instantaneous battery efficiency  $\eta_{battery}$ is defined as follow:

$$\eta_{battery} = \frac{|P_{stack}|}{|P_{stack}| + P_{mech}} \qquad [-] \tag{36}$$

Clearly, the battery performance is poor as it can be observed in Fig. 14 where  $\eta_{battery}$  is illustrated as a function of the stack current  $I_{stack}$  and the state of charge *SoC*. Indeed, the battery often consumes more power than necessary; therefore, constantly operating the battery at  $Q_{max}$  is not a wise strategy. Nevertheless, it is possible to improve this efficiency by limiting the operating range of the battery (smaller current and/or narrower state of charge); thus the flowrate  $Q_{max}$  and the mechanical power  $P_{mech}$  are reduced. But this also reduces the power rating and/or the energetic capacity while it increases the cost.

#### 7.2 Minimal flowrate

The low efficiency at constant flowrate  $Q_{max}$  is due to the large mechanical losses  $P_{mech}$ ; therefore, a second control strategy is proposed to minimize  $P_{mech}$ . In that case, the battery is operating at a minimal flowrate  $Q_{min}$  that is constantly adapted to the actual operating conditions (*SoC* and  $I_{stack}$ ) in order to supply just enough electroactive materials to fuel the electrochemical reactions. Since the vanadium concentrations  $c_V$  change proportionally to  $I_{stack}$ , there are critical operating points where  $c_V$  is close to its boundary. In some cases, the variations of vanadium concentrations tend toward the limit values (Fig. 13). In these critical regions, the electrolyte flowrate Q must be larger to palliate the scarcity of electroactive vanadium ions.

Hence, the minimal flowrate  $Q_{min}$  depends on the required amount of electroactive species, and in consequence on  $I_{stack}$ , and on the input vanadium concentrations  $c_{in}$  that are either

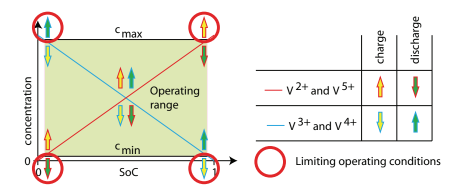


Fig. 13. Operating range and limiting operating conditions. The arrows represent the direction of the vanadium concentrations change as a function of the battery operating mode. The critical operating regions are highlighted in red; they represent the regions where the vanadium concentration  $c_{vanadium}$  tends to its limiting concentrations ( $c_{max}$  or  $c_{min}$ ).

being depleted ( $\downarrow$ ) or augmented ( $\uparrow$ ).  $Q_{min}$  can be derived from (17):

$$Q_{min,\downarrow}(t) = \frac{bN_{cell}i(t)}{F(c_{out,min} - c_{in,\downarrow}(t))} \qquad [l/s]$$
(37)

$$Q_{min,\uparrow}(t) = \frac{bN_{cell}i(t)}{F(c_{out,max} - c_{in,\uparrow}(t))} \qquad [l/s]$$
(38)

where  $c_{out,min}$  and  $c_{out,max}$  are constant minimal and maximal output concentrations. The limiting species depends on the operating mode (charge or discharge); thus  $Q_{min}$  is given by the maximal value of (37) and (38):

$$Q_{min}(t) = \max\left(Q_{min,\downarrow}(t), Q_{min,\uparrow}(t)\right) \qquad [l/s] \tag{39}$$

 $Q_{min}$  is illustrated in Fig. 14 for a wide spectrum of operating points; clearly,  $Q_{min}$  is larger in the critical regions that were highlighted in Fig. 13. Moreover,  $Q_{min}$  is, in comparison, very small in the other operating regions; therefore, there must be a large benefit to operate the battery at  $Q_{min}$ .

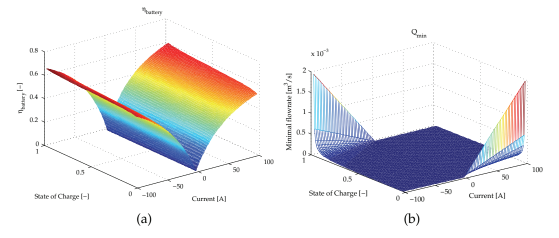


Fig. 14. (a) The battery efficiency  $\eta_{battery}$  at constant flowrate  $Q_{max}$  as a function of the state of charge *SoC* and current *I*. (b) Minimal flowrate  $Q_{min}$  as a function of the stack current  $I_{stack}$  and the state of charge *SoC*.

But a change in the flowrate Q also modifies the vanadium concentrations  $c_{cells}$  within the cells according to (18), and in consequence the stack voltage  $U_{stack}$  and power  $P_{stack}$  according to (2) and (3). This phenomenon is illustrated in Fig. 15 where the equilibrium voltage E at  $Q_{max}$  and  $Q_{min}$  is shown: an increase of the flowrate has always a beneficial effect on E. Furthermore, the equivalent state of charge  $SoC_{eq}$  which represents the SoC of the electrolyte within the cells is also illustrated as a function of Q. Clearly,  $SoC_{eq}$  tends toward the battery SoC at high Q. Therefore, the change in  $c_{cells}$  is maximal at  $Q_{min}$ ; and consequently a large variation of  $U_{stack}$  and  $P_{stack}$  is expected between the operations at  $Q_{min}$  and  $Q_{max}$  as it can be observed in Fig. 16. From the strict point of view of  $P_{stack}$ , it is more interesting to operate the battery at  $Q_{max}$ ; indeed, more power is delivered during the discharge and less is consumed during the charge. But it will be shown in the next sections that the mechanical power greatly deteriorates the performance and that the energy efficiency at  $Q_{max}$  is unacceptable.

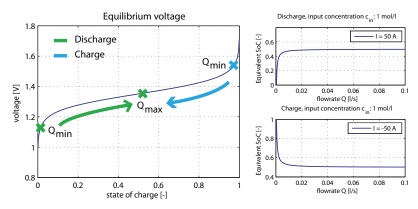


Fig. 15. Effect of the flowrate *Q* on the equilibrium voltage *E*. On the right, the variation of the equivalent state of charge *SoC* as a function of *Q* during the discharge and the charge. In this example, the battery *SoC* is 0.5, i.e. the input concentrations are 1 M for each vanadium species.

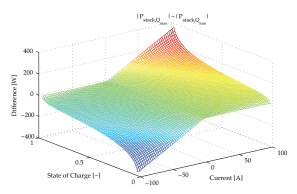


Fig. 16. The difference between the stack power  $|P_{stack,Qmax}|$  at  $Q_{max}$  and the stack power  $|P_{stack,Qmin}|$  at  $Q_{min}$ .

# 8. Optimal operating point at constant current

In the previous sections, the advantages and disadvantages of operating the battery at either  $Q_{max}$  and  $Q_{min}$  were discussed. At  $Q_{max}$ , the stack power  $P_{stack}$  has the highest possible value but the mechanical power  $P_{mech}$  is also very large and consequently deteriorates the performance. At  $Q_{min}$ ,  $P_{mech}$  is reduced to the minimum, but  $P_{stack}$  is negatively affected.

Therefore, it should exist an optimal flowrate  $Q_{opt}$  somewhere between  $Q_{min}$  and  $Q_{max}$  that increases  $P_{stack}$  while maintaining  $P_{mech}$  at a small value.

#### 8.1 Optimal flowrate during the discharge

In this section, the battery is controlled by the reference current  $I_{stack,ref}$ ; therefore there is only one control variable: the flowrate Q. Indeed, the stack power  $P_{stack}$  depends on  $I_{stack}$ , Qand the state of charge SoC whereas the mechanical power depends on Q and the electrolyte properties: the density  $\rho$  and the viscosity  $\mu$  that are maintained constant in this work. During the discharge, the optimal operating point is found when the flowrate  $Q_{opt}$  maximizes the power delivered by the stack  $P_{stack}$  while minimizing the mechanical power  $P_{mech}$ . When these conditions are met together, the power delivered by the battery  $P_{VRB}$  is optimized:

$$\max(\underbrace{P_{VRB}}_{f(U_{stack}, I_{VRB})}) = \max(\underbrace{P_{stack}}_{f(I_{stack}, Q, SoC)} - \underbrace{P_{mech}}_{f(Q, \mu, \rho)})$$
(40)

In Fig. 17,  $P_{VRB}$  is represented during the discharge as a function of Q at different states of charge for a current of 100 A. Clearly, an optimal flowrate  $Q_{opt}$  exists between  $Q_{min}$  and  $Q_{max}$  that maximizes  $P_{VRB}$ . The shape of the curves can be generalized to other discharge currents  $I_{stack} > 0$ ; although in some cases where  $I_{stack}$  is low,  $P_{VRB}$  might become negative at inappropriately high flowrate Q.

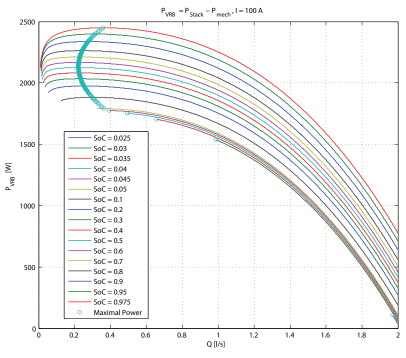


Fig. 17. Optimal flowrate  $Q_{opt}$  as a function of the flowrate Q and the state of charge *SoC*. Note that when *SoC* is low,  $Q_{opt}$  is equal to the minimal flowrate Q, and the discharge current is equal to 100 A.

#### 8.2 Optimal flowrate during the charge

At constant current  $I_{stack,ref}$ , the quantity of electrons  $e^-$  stored in the electrolyte does not depend on the stack power  $P_{stack}$  but solely on the stack current  $I_{stack}$ ; therefore, there is no reason to have a high  $P_{stack}$ . Hence, the optimal flowrate  $Q_{opt}$  during the charge is found

when the sum of  $P_{stack}$  and  $P_{mech}$  is simultaneously minimal. This condition is expressed by the following relation<sup>3</sup>:

$$\min(\underbrace{|P_{VRB}|}_{f(U_{stack}, I_{VRB})}) = \min(\underbrace{|P_{stack}|}_{f(I_{stack}, Q, SoC)} + \underbrace{P_{mech}}_{f(Q, \mu, \rho)})$$
(41)

The optimal flowrate  $Q_{opt}$  is illustrated in Fig. 18 where  $P_{VRB}$  is shown as a function of Q and *SoC*. At very high *SoC*,  $Q_{opt}$  is equal to  $Q_{min}$  because the electrolyte carries a very small amount of electroactive vanadium ions. Again, the shape of the curves can be generalized to other charge currents  $I_{stack} < 0$ .

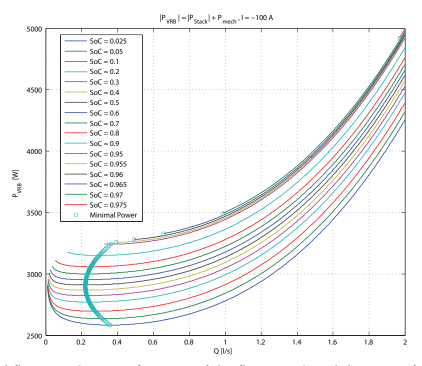


Fig. 18. Optimal flowrate  $Q_{opt}$  as a function of the flowrate Q and the state of charge *SoC*. Note that when *SoC* is high,  $Q_{opt}$  is equal to the minimal flowrate Q, and that the charge current  $I_{stack}$  is equal to -100 A.

# 8.3 Charge and discharge cycles

It is always difficult to assess the performance of a battery because it often depends on the operating conditions. In this section, a series of charge and discharge at constant current is performed at minimal flowrate  $Q_{min}$ , at maximal flowrate  $Q_{max}$  and at optimal flowrate  $Q_{opt}$  in order to assess the performance of this new control strategy.

The voltage  $\eta_{voltage}$  and energy  $\eta_{energy}$  efficiencies are summarized in Tab. 8 and 9; the coulombic efficiency  $\eta_{coulombic}$  is in all cases equal to 100% because the model does not take into account any side reactions such as oxygen or hydrogen evolution nor any cross mixing of the electrolyte.

Both  $\eta_{voltage}$  and  $\eta_{energy}$  decrease when the current increase; this is mainly due to the internal losses  $U_{losses}$  that are proportional to the current  $I_{stack}$ , although the flowrates  $Q_{min}$  and  $Q_{opt}$  increases to supply enough electroactive species. The highest voltage efficiencies occur

 $<sup>^{3}</sup>$ A close look at this relation reveals that it is the same as (40), but (41) is more intuitive for the charge.

Current [A]	η <sub>voltage,Qmax</sub> [%]	η <sub>voltage,Qmin</sub> [%]	η <sub>voltage,Qopt</sub> [%]
10	97.02	87.74	96.69
20	94.13	85.11	93.70
40	88.58	80.09	88.03
60	83.33	75.31	82.69
80	78.37	70.78	77.68
100	73.65	66.46	72.94

Table 8. Stack voltage efficiency  $\eta_{voltage}$  at constant maximal flowrate  $Q_{max}$ , at minimal flowrate  $Q_{min}$  and at optimal flowrate  $Q_{opt}$ .

Current [A]	Time [h]	η <sub>energy,Qmax</sub> [%]	η <sub>energy,Qmin</sub> [%]	η <sub>energy,Qopt</sub> [%]
10	44.49	-73.42	87.73	96.54
20	22.24	-53.34	85.10	93.51
40	11.12	-25.65	80.04	87.77
60	7.41	-8.17	75.31	82.34
80	5.56	3.24	70.78	77.26
100	4.45	10.81	66.24	72.43

Table 9. Overall VRB energy efficiencies  $\eta_{energy}$  at constant maximal flowrate  $Q_{max}$ , at minimal flowrate  $Q_{min}$  and at optimal flowrate  $Q_{opt}$ .

at  $Q_{max}$  because of its positive effect on the stack voltage  $U_{stack}$  highlighted in section 7.2; consequently, the worst voltage efficiencies occur at  $Q_{min}$ . Moreover, the voltage efficiencies at  $Q_{opt}$  are very close to the maximal efficiencies obtained at  $Q_{max}$ . In fact, the stack voltages  $U_{stack,Qmax}$  and  $U_{stack,Qopt}$  are very close as it can be observed in Fig. 19.

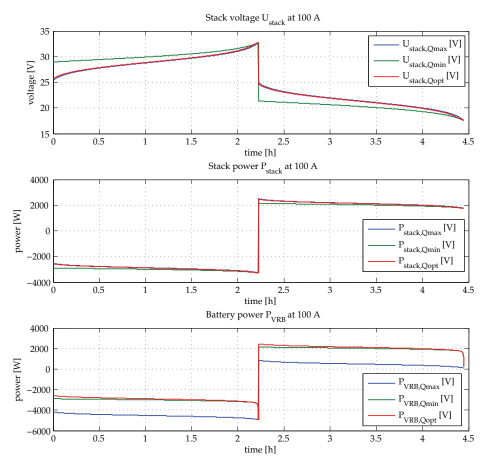


Fig. 19. Stack voltage  $U_{stack}$ , stack power  $P_{stack}$  and battery power  $P_{VRB}$  during a charge and discharge cycle at 100 A.

Obviously, operating the battery at  $Q_{max}$  is a problematic strategy as  $\eta_{energy,Qmax}$  is very small or even negative: at small currents, the battery does not deliver any power to the load but consumes more power to operate the pumps than the stack is furnishing. When  $P_{mech}$  is minimized, the energy efficiencies already become interesting at  $Q_{min}$ , but they are increased by a further 10% when the battery is operating at  $Q_{opt}$ .

In order to compare the model with experimental data, the stack characteristics were defined to match the stack presented in section 3.7. The experimental results of M. Skyllas-Kazacos and al. are summarized in Tab. 4 (Skyllas-Kazacos & Menictas, 1997); note that they do not take into account the mechanical power required to operate the pumps and that the flowrate was constant (2 l/s which correspond to  $Q_{max}$ ). The losses in coulombic efficiency  $\eta_{coulombic}$  can be caused by side reactions or cross mixing of electrolyte through the membrane which are not taken into account in the model; but  $\eta_{coulombic}$  improves as the battery becomes conditioned. In that case, the energy efficiency  $\eta_{energy,Qopt}$  at optimal flowrate is very close to the maximal electrochemical energy efficiency. Finally, a very good concordance is observed between the voltage efficiencies at  $Q_{max}$  and the experimental results.

# 9. Optimal operating point at constant power

In practice, the battery must often deliver a certain amount of power to the load: the battery is controlled by a reference power  $P_{ref}$ . In that case, a second control variable is available in supplement of the flowrate Q: the stack current  $I_{stack}$ . The optimal operating point is the couple  $Q_{opt}$  and  $I_{opt}$  that maximizes the amount of charge that are stored within the electrolyte during the charge and minimizes the amount of charge that are consumed during the discharge. These conditions can be related to  $I_{stack}$ :

$$\underbrace{P_{VRB}}_{constant} = \underbrace{P_{stack}}_{f(I_{stack},Q,SoC)} - \underbrace{P_{mech}}_{f(Q,\mu,\rho)} [W]$$
(42)

during the charge: 
$$\max(|I_{stack}|)$$
 [A] (43)

during the discharge: 
$$min(I_{stack})$$
 [A]

Again, an optimal operating point exists in between the maximal  $Q_{max}$  and minimal  $Q_{min}$  flowrates as it can be observed in Fig. 20 where operating points are represented for different battery power  $P_{VRB}$  during the discharge at a *SoC* equal to 0.5. At the optimal flowrate  $Q_{opt}$ , the battery delivers the same power  $P_{VRB}$  but consumes less active vanadium ions; therefore, the battery will operate longer and deliver more power.  $Q_{opt}$  increases with  $P_{VRB}$  until it reaches a plateau due to the transition between the laminar and the turbulent regime.

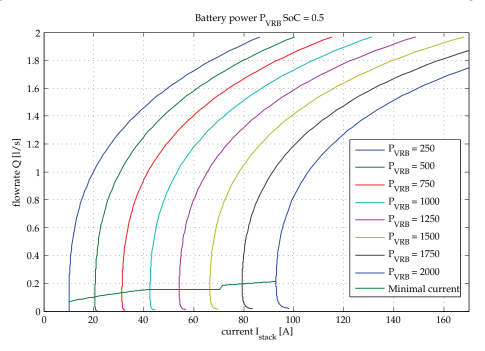


Fig. 20. Battery power  $P_{VRB}$  as a function of the discharge current  $I_{stack}$  and the electrolyte flowrate Q at a state of charge *SoC* equal to 0.5. The optimal operating points occurs when the current  $I_{stack}$  is minimal for a given battery power  $P_{VRB}$ .

In fact,  $I_{stack}$  increases above the optimal flowrate to compensate the higher mechanical loss: the stack must deliver more power. Below  $Q_{opt}$ ,  $I_{stack}$  increases this time to compensate the lower stack voltage  $U_{stack}$  due to the lower concentrations of active species. The shape of the curves can be generalized for other states of charge *SoC*.

The optimal operating points during the charge are illustrated in Fig. 21 where the battery power  $P_{VRB}$  is shown as a function of the current  $I_{stack}$  and the flowrate Q at a state of charge

of 0.5. The optimal operating point maximizes the current  $|I_{stack}|$  delivered to the stack in order to store the maximum amount of electroactive species at a given power  $P_{VRB,ref}$ ; again, the optimal flowrate  $Q_{opt}$  increases with the battery power  $P_{VRB}$  until it reaches the plateau due to the flow regime transition.

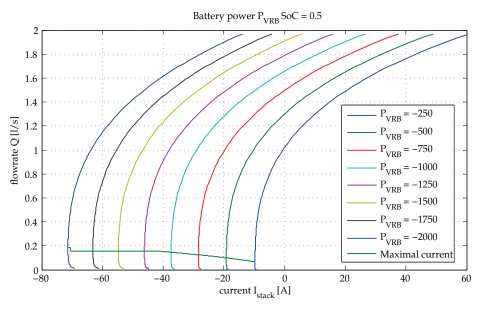


Fig. 21. Battery power  $P_{VRB}$  as a function of the charge current  $I_{stack}$  and the electrolyte flowrate Q at a state of charge *SoC* equal to 0.5. The optimal operating points occurs when the current  $|I_{stack}|$  is maximal for a given battery power  $P_{VRB}$ .

Interestingly, we observe in Fig. 21 that the stack current  $I_{stack}$  changes its sign at high flowrate Q; in these unacceptable conditions, the stack is discharged while the battery is being charged. During the charge, the stack current  $|I_{stack}|$  decreases above the optimal flowrate  $Q_{opt}$  to compensate the higher mechanical loss  $P_{mech}$ ; in consequence, less power is available to charge the stack (see (42)). Below the optimal flowrate  $Q_{opt}$ , the stack current  $|I_{stack}|$  also decreases because the stack voltage  $U_{stack}$  increases due the change in electroactive species concentrations within the cells  $c_{cell}$ ; note that the mechanical power  $P_{mech}$  is also reduced below  $Q_{opt}$ . Furthermore, the shape of the curves in Fig 21 might be generalized to other states of charge SoC.

# 9.1 Charge and discharge cycles

A new series of charge and discharge cycles at constant power was performed to determine the energy efficiencies at minimal flowrate  $Q_{min}$  and at the optimal operating point:  $I_{opt}$  and  $Q_{opt}$ . This optimal point is constantly determined as a function of the actual conditions. The energy efficiencies are given in Tab. 10. The energy efficiency at optimal flowrate  $\eta_{energy,Qopt}$  is increased by 10% at maximal power when compared to battery operations at minimal flowrate  $Q_{min}$ .

# 10. Epilogue

Today, the electricity industries are facing new challenges as the market is being liberalized and deregulated in many countries. Unquestionably, electricity storage will play, in the near future, a major role in the fast developing distributed generations network as it has

Power [W]	η <sub>energy,Qmin</sub> [%]	η <sub>energy,Qopt</sub> [%]
500	87.07	93.59
1000	81.04	87.78
1500	75.47	82.09
2000	69.91	76.39
2500	63.97	70.48

Table 10. Overall VRB energy efficiencies  $\eta_{energy}$  for a charge and discharge cycle at constant power at either optimal flowrate  $Q_{opt}$  and minimal flowrate  $Q_{min}$ .

many advantages to offer: management of the supply and demand of electricity, power quality, integration of renewable sources, improvement of the level of use of the transport and distribution network, etc. Over the years, many storage technologies have been investigated and developed, some have reached the demonstrator level and only a few have become commercially available. The pumped hydro facilities have been successfully storing electricity for more than a century; but today, appropriate locations are seldom found. Electrochemical storage is also an effective means to accumulate electrical energy; among the emerging technologies, the flow batteries are excellent candidates for large stationary storage applications where the vanadium redox flow battery (VRB) distinguishes itself thanks to its competitive cost and simplicity.

But a successful electricity storage technology must combine at least three characteristics to have a chance to be widely accepted by the electrical industry: low cost, high reliability and good efficiency. A lot of works have already been done to improve the electrochemistry of the VRB and to reduce its overall manufacturing cost. With the multiphysics model proposed in this chapter, we are able to address primarily the battery performance and indirectly its cost; indeed, a good efficiency enhances the profitability and consequently reduces the operating cost.

This ambitious model encompasses the domains of electricity, electrochemistry and fluid mechanics, it describes the principles and relations that govern the behaviour of the VRB under any set of operating conditions. Furthermore, this multiphysics model is a powerful means to identify and quantify the sources of losses within the VRB storage system; indeed, we are now able to understand how the VRB operates and to propose strategies of control and operation for a greater effectiveness of the overall storage system.

Another important feature of this multiphysics model is to facilitate the integration of the VRB into the electrical networks. Indeed, power converters, whose properties and characteristics are known and efficient, are required in practice to interface the VRB with the network; the overall performance might improve if their control strategy takes into account the VRB characteristics.

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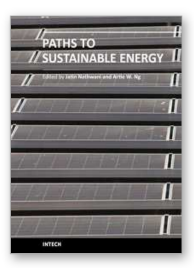
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The world's reliance on existing sources of energy and their associated detrimental impacts on the environment- whether related to poor air or water quality or scarcity, impacts on sensitive ecosystems and forests and land use - have been well documented and articulated over the last three decades. What is needed by the world is a set of credible energy solutions that would lead us to a balance between economic growth and a sustainable environment. This book provides an open platform to establish and share knowledge developed by scholars, scientists and engineers from all over the world about various viable paths to a future of sustainable energy. It has collected a number of intellectually stimulating articles that address issues ranging from public policy formulation to technological innovations for enhancing the development of sustainable energy systems. It will appeal to stakeholders seeking guidance to pursue the paths to sustainable energy.

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