Electrolysis

In 1800, the two English scientists William Nicholson and Anthony Carlisle discovered direct current electrolysis, thereby establishing electrochemistry as a new scientific field. For a long time, electrolysis was the dominant technology for the industrial generation of hydrogen. However, the increasing use of natural gas largely displaced this technology with the more cost-effective steam reforming.

Now, over 200 years later, the splitting of water with electrical current is experiencing a renaissance. This is because this technology provides the possibility of converting and storing regenerative electric power in the form of hydrogen as a chemical energy source. In this process, traditional alkaline electrolysis is supplemented by the modern technology of polymer electrolyte membrane (PEM) electrolysis. This is based on a publication by Russell et al. in 1973.¹ The high reaction rate and the high power density make PEM electrolysis interesting with the use of fluctuating renewable energy.²

The efficiency of the electrolysis systems is critical both technically and economically for electrochemical hydrogen generation using renewable energy. Hydrogen generation costs are primarily dominated by the relatively high power costs in addition to operating hours and amortization of the plant. From a technical standpoint, these costs can only be reduced by increasing overall efficiency.

The following document is intended to illustrate for the reader what has to be taken into account when specifying or comparing efficiency levels for electrolysis systems.
The efficiency of an electrolysis plant is one of its key parameters. It is largely determining for power consumption and therefore the bulk of operating costs. On the other hand, efficiency is interrelated with all of the peripheral parameters that also influence hydrogen generation costs, such as operating mode (operating hours) service life and investment costs. For that reason alone, the definition, calculation and interpretation of efficiency in electrolysis technology merit a closer look.

Stack, system and plant efficiency are often referred to in actual use of this technology. All three of these represent the products of various sub-efficiencies and are therefore not clearly defined. Stack efficiency, which is sometimes also known as DC efficiency, is itself the product of Faraday efficiency (also known as current efficiency) and the voltage efficiency, which is often used as a characteristic curve for an electrolysis process due to the dominating losses. In an ideal cell, voltage defines the efficiency over the range of current densities.

The so-called U/I characteristic is therefore also the most important graph in the evaluation of electrolysis technology. In addition to the dominating losses in the cell, various auxiliary loads must be accounted for in determining the efficiency of a system or plant. The greatest challenge in this regard is to clearly define the boundaries of the system or balance. This affects not only the individual packages and their power consumption, but also the condition and quality of all parameters at the balance interfaces, such as water quality, waste heat, the quality of the electrical power and gas quality and pressure level.

![Figure 1: Schematic representation of a U/I characteristic and its indirectly proportional effect on efficiency.](image)
Electrolysis efficiency

Definitions of efficiency

A systematic examination of the efficiency of an electrolysis plant for hydrogen generation yields many specifications for a wide range of system boundaries. The most frequently used parameters are presented below.

**Faraday efficiency**

Faraday efficiency is the ratio of the actually produced and technically usable volume of hydrogen production and the theoretical maximum production volume. For example, it accounts for $H_2$ diffusion losses through the membrane, electrical current losses or hydrogen losses through the conversion of oxygen contamination in the downstream hydrogen purification system. Mass losses due to sampling or other losses can also be included in the calculation.

$$\varepsilon_F = \frac{m_{\text{re}H_2}}{m_{\text{id}H_2}}$$  \hspace{1cm} (8)

$m_{\text{re}H_2}$ is defined as the actual production volume of usable hydrogen and $m_{\text{id}H_2}$ as the maximum possible hydrogen production volume under ideal conditions.

**Voltage efficiency**

Voltage efficiency is often confused with DC efficiency. Voltage efficiency is defined as the quotient of thermoneutral voltage $E_{th}$ and the measured cell voltage $E_{\text{cell}}(i,T)$. Thermoneutral voltage comprises the heat energy bound by the change in entropy and is $1.48$ V under standard conditions. This is not to be confused with the reversible cell voltage ($E_{rev} = 1.23$ V) that must be applied as a minimum to split water into its components of $H_2$ and $O_2$. The cell voltage, however, also includes all of the catalytic and electrical losses within the DC-powered electrochemical cell.

$$\varepsilon_V = \frac{E_{th}}{E_{\text{cell}}(i,T)}$$  \hspace{1cm} (9)

**DC efficiency**

DC efficiency comprises the efficiency of the stack or of a module. Ultimately, it represents the ratio of the energy input to the stack in the form of DC current and the energy content of the technically usable hydrogen at the stack output. Specification of the higher or lower heating value is relevant here.

$$\varepsilon_{DC} = \frac{V_{H_2}H_2}{P_{elDC}}$$  \hspace{1cm} (10)

The product with voltage efficiency can also be used depending on the definition of the system boundaries for Faraday efficiency.

$$\varepsilon_{DC} = \varepsilon_V \cdot \varepsilon_F$$  \hspace{1cm} (11)

**System/plant efficiency**

While the hydrogen community is somewhat in agreement regarding the above efficiency definitions, with the exception of the nomenclature, the system boundaries for system or plant efficiency are defined arbitrarily. Depending on the scope of supply, losses may also be included from AC/DC conversion and transformation from medium voltage, water treatment, the cooling systems, the provision of electrical energy for buildings and auxiliary packages or compression and hydrogen purification.

The respective system boundaries must be investigated in corresponding detail to enable the comparison of efficiency data.

$$\varepsilon_{SyS} = \frac{V_{\text{interface customer}}H_2}{P_{elAC\text{interface grid}}$$  \hspace{1cm} (12)
U/I characteristic

As for most technical equipment, efficiency also depends on the operating point. The delightful aspect of electrochemistry is that the efficiency of the electrochemical reaction, i.e. voltage efficiency, increases at part load. Temperature also has a significant effect on efficiency, as does pressure to a lesser extent.

Operating parameter I – current density

An electrolysis power curve is typically represented as a U/I characteristic.

Operating parameter II – temperature

Temperature has a further significant effect on efficiency. Both the catalytic reaction rate as well as specific resistance are heavily dependent on temperature. The higher the temperature, the higher the efficiency.

Why not simply operate at low current densities?

The higher the current density, the higher the rate of hydrogen production. This means that the specific costs for an electrolyzer increase significantly at lower current density. The correct current density should always be optimized with the objective of minimizing hydrogen production costs. The objective is to obtain an optimum balance between high efficiency and service life.
Increasing voltage efficiency
The slope of the U/I characteristic is thus a true indicator for efficiency improvement. A smaller slope can be achieved with more effective catalysts or lower resistances in the cell.

![Figure 6: Schematic showing the effect of lower resistance on efficiency](image)

The distance between the two electrodes has a large effect on the slope of the U/I characteristic. The shorter the distance the charge carriers have to traverse between the anode and the cathode, the smaller the voltage drop is, with correspondingly higher efficiency.

**Why not just move the electrodes closer together?**
For safety reasons alone, the oxygen generated at the anode shall not be allowed to mix with the hydrogen generated at the cathode. The smaller the distance, the higher the diffusion rate of contaminant gases from one side to the other if the material remains the same. This results in higher Faraday losses and to higher contamination of the product gases.

**PEM and alkaline electrolysis – specific differences**
There are currently two commercially relevant electrolysis technologies on the market that are implemented for production in the MW range: PEM and alkaline electrolysis. Both technologies differ primarily in the ionic charge carrier that electrically closes the electrochemical process, as well as in the actual electrolytes.

Polymerelectrolyte membrane (PEM) or proton exchange membrane technology, as the name indicates, includes a solid electrolyte that electronically isolates the anode from the cathode but electrically closes the electric circuit through its selective conductivity for cations. This nearly gas-tight solid electrolyte offers several specific advantages. On the one hand, it serves as a physical divider between the anode (oxygen side) and the cathode (hydrogen side) and prevents mixing of the generated gases. This enables operation with a differential pressure. Unintentional pressure differentials that could result in mixing of the gases and thus to a safety hazard can therefore be more easily handled. On the other hand, the membrane also ensures a high gas product purity in dynamic operation or during more extended part-load operation where the contaminant gas concentration in both electrolysis technologies increases specifically as a function of the gas production rate due to diffusion.

![Figure 7: Schematic of PEM and alkaline electrolyzers](image)

Alkaline electrolyzer have a porous separator to physically separate the hydrogen and oxygen gases while still enabling transport of the liquid electrolyte. This porous separator increases the requirements for alkaline electrolyzers in dynamic operation. Differential pressures that can result in hazardous mixing of oxygen and hydrogen absolutely must be prevented in the alkaline electrolyzer due to its porous separator.
Differences in characteristic
The electrodes for alkaline electrolysis are traditionally further apart from each other than those in PEM electrolysis. In addition to different electrode materials, this generally results in a steeper U/I characteristic for alkaline electrolyzers. The conductivity of the electrolytes also plays a significant role.

In addition to ionic conductivity, a 30% KOH solution also has a considerably higher electrical conductivity than that of deionized water. This results in so-called shunt currents (loss currents), i.e. electrons that do not participate in the electrochemical reaction but rather flow through the electrolyte to the adjacent cell. These must be prevented or assigned to Faraday efficiency.

The question of comparing the efficiency of both technologies is therefore not easy to answer. The fact is that the selection of the operating point and the associated voltage are decisive for efficiency. The generally low overall resistances in PEM technology result in a broader range of operating modes. While PEM electrolysis is operated at between 0 and 3 A cm$^{-2}$, alkaline electrolyzers permit significantly lower current densities to achieve a comparable voltage and therefore also a comparable efficiency.

Further fundamental principles 2,3
Efficiency $\varepsilon$ is a technical parameter that is generally specified with values of between 0 and 1 or 0% and 100%. In principal, the definition of efficiency is generally applicable and can be described with the following equation:

$$\varepsilon = \frac{\text{Useful}}{\text{Expended}}$$  \hspace{1cm} (1)

The term efficiency is most frequently found in connection with energy flow $E$. Useful energy is in the numerator and the energy expended is in the denominator. Power $P$ can also be used if no energy is stored within the balance boundaries.

Efficiency of an ideal galvanic cell
The attractive aspect of electrochemistry is the direct conversion of chemical energy to electrical energy and vice versa. This advantage can be illustrated by a comparison of the maximum theoretical efficiency of a thermal power plant with that of a galvanic cell such as a fuel cell. Instead of converting chemical energy first into thermal energy, then into mechanical energy and finally into electrical energy, electrochemistry accomplishes this in a single step.

Efficiency of an ideal galvanic cell
$$\varepsilon = \frac{E_{\text{Useful}}}{E_{\text{Expended}}} = \frac{P_{\text{Useful}}}{P_{\text{Expended}}}$$  \hspace{1cm} (2)

The description of the classic Carnot cycle defines the maximum theoretical efficiency with the following formula:

$$\varepsilon_C = 1 - \frac{T_u}{T}$$  \hspace{1cm} (3)

Where $T$ is the maximum process temperature and $T_u$ the ambient temperature. The higher the temperature of the Carnot cycle, the higher the efficiency of the thermal power plant.

The ideal efficiency $\varepsilon_{id}$ of a galvanic cell comprises the quotient of Gibbs free energy $\Delta G$ and overall reaction enthalpy $\Delta H$.

$$\varepsilon_{id} = \frac{\Delta G}{\Delta H} = 1 - \frac{T_u}{T}$$  \hspace{1cm} (4)

In comparison with the ideal efficiency of a classic thermal power plant, the theoretical efficiency of the galvanic cell is thus significantly higher over a wide temperature range.

Figure 8: Operating range for alkaline and PEM electrolyzers

Figure 9: Comparison of the energy conversion chain between heat engine and fuel cell

Figure 10: Ideal efficiency curves for a galvanic cell and for the Carnot cycle
Table 1: Example calculation for various standard conditions at the same percentage efficiency

<table>
<thead>
<tr>
<th>Standard</th>
<th>Condition</th>
<th>Density in kg/m³stp</th>
<th>Efficiency in kWh/m³stp</th>
<th>Efficiency in %</th>
</tr>
</thead>
<tbody>
<tr>
<td>DIN 1343</td>
<td>$p_n = 1.01325$ bar; $T_n = 273.15$ K</td>
<td>0.0899</td>
<td>4.425</td>
<td>80</td>
</tr>
<tr>
<td>Linde/Air Liquide</td>
<td>$p_n = 1$ bar; $T_n = 288.15$ K</td>
<td>0.0841</td>
<td>4.1375</td>
<td>80</td>
</tr>
<tr>
<td>ISO 2533</td>
<td>$p_n = 1.01325$ bar; $T_n = 288.15$ K</td>
<td>0.0852</td>
<td>4.158</td>
<td>80</td>
</tr>
</tbody>
</table>

Upper or lower heating value
The expenditure associated with a forced cycle such hydro electrolysis is the electrical power that must be supplied to the system. Useful output from the electrolysis of water is, however, a matter of definition. If the hydrogen used after generation is reused for energy production after it is generated (conversion to mechanical, electrical or thermal energy), the lower heating value $H_u$ (LHV) must be used under consideration of the entire conversion chain.

$$\varepsilon_{LHV} = \frac{\dot{V}_H \cdot H_u}{P_{\text{electrical}}}(5)$$

However, if instead of the entire conversion chain, only the hydrogen is regarded as a chemical product for use in the chemical industry or as a technical gas, the fuel value or higher heating value $H_o$ (HHV) can be used for calculating efficiency. The necessary reaction enthalpy for converting the liquid reactant water to gaseous hydrogen corresponds to the higher heating value.

$$\varepsilon_{HHV} = \frac{\dot{V}_H \cdot H_o}{P_{\text{electrical}}}(6)$$

In general, the higher heating value is used when referring to the electrolysis of water. This is due to the fact that the use to which the hydrogen is put should not be assumed by the manufacturer and the pure percentage value based on the higher heating value suggests a significantly higher efficiency than that based on the lower heating value.

Deviation from specified percentage efficiency
As an alternative to the typical specification of efficiency as a percentage, the efficiency of hydrogen electrolysis can also be communicated in a mixed form between the energy and product stream. The following values are often used here:

$$\frac{\text{kWh}}{\text{m³stp}} \text{ or } \frac{\text{kWh}}{\text{kg}} (7)$$

While the representation of efficiency as a percentage requires that the reference value of higher or lower heating value also be indicated, standard conditions must be indicated when communicating energy expenditure per unit volume of gas. There is more than just one definition of the "standard conditions" of a volume, which significantly affects the energy content due to differences in the reference temperature and pressure. The specification of energy expenditure per unit volume is therefore ambiguous without also specifying the standard conditions.

Conclusion
It must be ensured that all relevant reference points are given when evaluating the efficiency of electrolysis systems. This starts with the definition of the system boundaries and standard conditions, affects part-load behavior and dynamic conditions and ends, not least, with degradation. All of these factors ultimately have an effect on efficiency.

A system manufacturer must strike an optimal balance to enable the lowest hydrogen generation costs while providing maximum availability for the customer over the long term. On the other hand, it is the task of interested parties, be they a customer or the author of a study, to define all of the required reference points to enable an objective comparison.
Sources


Index of Abbreviations

PEM  Polymer electrolyte membrane
DC   Direct current
AC   Alternating current

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