Benchmark Synthetic Training Data for Artificial Intelligence-based Li-ion Diagnosis and Prognosis

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Abstract: Accurate lithium battery diagnosis and prognosis is critical to increase penetration of electric vehicles and grid-tied storage systems. They are both complex due to the intricate, nonlinear, and path-dependent nature of battery degradation. Data-driven models are anticipated to play a significant role in the behavioral prediction of dynamical systems such as batteries. However, they are often limited by the amount of training data available. In this work, we generated the first big data comprehensive synthetic datasets to train diagnosis and prognosis algorithms. The proof-of-concept datasets are over three orders of magnitude larger than what is currently available in the literature. With benchmark datasets, results from different studies could be easily equated, and the performance of different algorithms can be compared, enhanced, and analyzed extensively. This will expend critical capabilities of current AI algorithms, tools, and techniques to predict scientific data.

Keywords: Machine Learning, Training Data, alawa, AI

In recent years, artificial intelligence (AI) has attracted a lot of attention for energy applications [1-4]. For the diagnosis and prognosis of lithium-ion battery (LiB), one bottleneck is the training data that is not populated enough and not representative of the projected sporadic usage [3,5]. This can hinder performance as models can only be as good as the data they were trained with. Most studies had training datasets below 20 samples. Among the studies with more, [5-12], the study by Severson et al. [5,11] stands out with 124 different conditions tested, although only charging conditions were varied. Even though online databases [5,13-18] are a step in the right direction, this is vastly insufficient because LiB degradation is path-dependent and small changes in conditions were shown to lead to drastic differences in durability [19]. This path dependence is an essential aspect to consider for the validation of online diagnosis and prognosis tools [20,21]. With a limited set of training data, the universality of the diagnosis and prognosis tool cannot be proven.

The need for high throughput computational data generation was highlighted in recent reviews [1,2], although no effort towards computational cycling training data has been reported. In this work, we will report the first synthetic benchmark training datasets that englobe the entire degradation spectrum. This will be done using computer-assisted voltage curve generation to remove the need for lengthy and costly experimental campaigns. The datasets will be computed using the mechanistic modeling approach we pioneered, along with other groups, in the mid-2000s [22-25]. The approach has been well validated [26-28] and has been intensively used in recent years with the rise in popularity of the electrochemical voltage spectroscopies (EVS) [29,30]. The use of the mechanistic approach will enable the creation of training datasets several orders of magnitude larger than the current ones encompassing all possible degradation scenarios. This is especially important for prognosis tools as battery capacity loss is known to accelerate at some point upon aging. This is the essential feature than any good prognosis needs to capture. This acceleration usually originates from degradations that have an “incubation” period that does not affect capacity [21]. A complete training dataset should contain examples of paths with an acceleration induced by different components of the degradation, whether it is the loss of lithium, losses in active materials, or resistance.
Compared to traditional battery modeling [31,32], the mechanistic framework is a backward modeling approach where the input is the degradation and the output the cell’s voltage and capacity. This makes the method perfect for generating training data and more efficient than electrochemical models because there is no need to find actual physical phenomena that could lead to a given degradation. The initial set up is also simpler without a need for complex parameterization and the only prerequisite being half-cell data versus a reference electrode for both electrodes. In 2017, we proposed to use this approach to perform sensibility analyses by simulating a wide range of hypothetical degradation scenarios [20]. This work is building on our previous effort to enable the creation of comprehensive training data.

In this work, we will focus on diagnosis and prognosis, two applications that require different sets of training data. To illustrate the difference between the two requirements, one can think of two time-dependent processes, one linear and one exponential, intersecting. At the intersection, the diagnostic will be the same, but the prognosis needs to be different. In addition, we will also address another hurdle for AI algorithms, the quest for meaningful learnable parameters. Although some studies seem to fit the full constant current voltage data [33], most studies favor the use of features of interest (FOI) and only focus on a specific part of the electrochemical response. This could be capacity and resistance evolution [10,34-37], curvature [38,39], sections of the voltage response [8,40-42], electrochemical impedance spectroscopy [43,44], variance [5,11], or EVS [33,45-48]. The latter has attracted a lot of attention in recent years since the early work on the technique [22,49,50]. Still, the correlation of the variations with degradation is not trivial and requires significant sensibility analysis to derive universal parameters [20]. Some FOIs proposed in the literature were proven not applicable outside of the tested data [21], which illustrates the need for more extensive training sets and proper sensitivity analysis of the learnable parameters. With controlled and complete training, the theoretical framework to establish relationships between data and models can be explored in much more detail, and results from different studies could be easily equated.

The purpose of this publication is to introduce a big data methodology and showcase its possibilities. The optimization of the technique to reduce computation time and its integration with AI algorithms is out of the scope of this work.

**Methodology**

The mechanistic approach conceptually follows the one described by Christensen and Newman to simulate cell degradation in the early 2000s [31,32]. Instead of computing-intensive electrochemical models to simulate half-cell electrode behavior, the mechanistic approach [22-25] uses experimental half-cell data for each electrode. Figure 1(a) presents the mechanistic representation of a cell. The matching consists of two parameters, the loading ratio LR between the capacities of both electrodes and their offset OFS that is representative of the amount of lithium lost during the solid electrolyte interphase formation. The circles in Figure 1(a) showcase how the experimental data overlap the emulated cell perfectly. To enhance the differences, the same data can be derived and plotted as incremental capacity, dQ/dV = f(V), Figure 1(b-d). For this EVS, each peak represents a phase transformation in one of the electrodes. A perfect match between the emulated and the experimental data proves that the emulated cell is perfectly replicating the experimental cell voltage variations. Figure 1(b-d) presents examples of the quality of the obtainable fits for cells based on different Li-ion chemistries from our previous work [51-53].
Figure 1: (a) Mechanistic model principles. The full cell response, in blue, corresponds to the response of the positive electrode, in red, minus the one of the negative electrode, in black, after matching. Fit for cells based on (b) a blended Graphite (Gr)-Silicon (Si) NE and Ni$_{0.8}$Mn$_{0.1}$Co$_{0.1}$O$_2$ (NMC811) PE [53], (Gr) negative electrodes (NE) and (c) Ni$_x$Al$_{1-x}$O$_2$ (NCA) [51] as well as a (d) blend of LiCoO$_2$ (LCO) and NCA as (PE) [52].

While the active materials are considered stable with aging, their quantity, as well as the amount of Lithium reacting, will change upon degradation. Degradation will then not affect the electrode OCV curves, but it will impact their matching. If less active material is available, the loading ratio between the electrodes will change. If some reactant is lost, the synchronicity of the electrodes will change. These matching changes can be rendered in the mechanistic approach via some scaling of the electrode curves and a translation of one electrode compared to the other, in other words changing LR and OFS in Figure 1(a). Figure 2(a) presents the impact of the variation of LR and OFS on cell capacity. It can be seen that there is an infinity of combination of LR and OFS that can lead to any capacity loss; this is the path dependence of the degradation. This also explains why measuring capacity will never be a good enough diagnosis of true degradation. Changes in the amount of active material are referred to as loss of active material (LAM). LAM can occur at the PE and NE. Change in the amount of lithium reacting is referred to as loss of lithium inventory (LLI). The thermodynamic degradation of Li-ion batteries can be decomposed into these three main degradation modes. LLI is inducing translation and the LAMs scaling, Figure 2(b-d). More details on the technique and the associated equations can be found in [25].
Figure 2: (a) Impact of changes of LR and OFS on the capacity loss. Arrows represent the impact of LLI and LAMs on LR and OFS. (b) Impact of LAMPE and (c) impact of LAMNE with the scaling of the affected electrode compared to the other. (d) Impact of LLI with the translation of the NE compared to the PE. In all cases, the full cell response, in blue, corresponds to the response of the positive electrode, in red, minus the one of the negative electrode, in black, after matching.

The validity of the predicted impact of LAM and LLI on the electrochemical behavior of experimental cells was verified by independent studies [26-28]. Most notably, Kassem et al. [26] confirmed that LLI does induce electrode slippage between the NE and Schmidt et al. [54] verified that LAM does influence the LR by assembling cells with different amounts of PE versus a constant NE. Moreover, over the past decade, numerous studies [29,30] successfully used this approach to diagnose the degradation of commercial cells. Figure 3(a,b) presents two examples of the fit of the voltage response of aged commercial cells from previous works [53,55]. Figure 3(c,d) presents the comparison of an experimental (from [53]) and a synthetic dataset constructed solely from half-cell data and simple equations that modified the matching of the electrodes based on the quantification of LLI and the LAMs. This highlights how the mechanistic approach is efficient to replicate the voltage variations associated with cell degradation.
Figure 3: Example of voltage emulation after aging for (a) a Gr//NCA discharge [55] after 1,500 equivalent full cycles based on electric vehicle driving and frequency regulation. (b) A Gr//LCO+NCA charges [53] after 1000 full cycles at 1.5C. (c) Experimental [53] and (d) synthetic voltage variations upon aging calculated from simple linear equations extracted from the diagnosis.

From the mechanistic approach point of view, each thermodynamic degradation can be described by a unique combination of the LLI, LAM PE, and LAM NE triplet. Conversely, scanning every possible combination of the triplet yield any potential degradation from which the voltage response can be reconstructed. This is the unique feature that is used in this work to build synthetic training datasets.

For datasets aimed at diagnosis, the approach used in previous work on an automated diagnosis tool for BMS [20] can be expanded. The [LLI, LAM PE, LAM NE] triplets can be normalized so that their sum is equal to one and they can then be represented in a ternary diagram, Figure 4. By scanning every point of this diagram (red dots) between 0 to 100% degradation for each mode (vertical lines), each possible degradation can be represented, and the associated voltage response calculated to be used in a training dataset. E.g., for the [0.5,0.2,0.3] triplet, LLI would be varied from 0 to 100% with 1% increments, LAM PE from 0 to 40% with 0.4% increments, and LAM NE from 0 to 60% with 0.6% increments so that their ratio is always conserved.
Figure 4: (a) Mechanistic model principles to emulate every possible degradation. Only partial degradation calculations were represented for illustration purposes.

For datasets aimed at prognosis, the evolution of each degradation mode needs to be calculated independently as the ratio between the normalized values of the triplet might not be constant. The triplet values could be calculated, per example, for different linear degradation paces, with paces linear combined with exponential variations, or with a delayed exponential increase, more representative of the real degradation of commercial cells [19,56,57]. Since the voltages associated with all the triplet values were already calculated for diagnosis, no additional simulations are actually necessary and the different degradation paths are all inscribed in Figure 4. They can be reconstructed by following the evolution of the triplet values in the ternary design space and the corresponding voltage variations can be accessed. Figure 5 presents an example of the process for two degradation paths for a Gr//LiFePO₄ (LFP) cell. Figure 5(a) plots the evolution of the triplets for each path. For each cycle, as exemplified by the black box in Figure 5(a), a triplet value can be determined and the quantification can be used to find the corresponding point in the ternary design space, Figure 5(b). In this example, path 2 (green) was simulated with linear variations for the three components of the triplet, its path is, therefore, vertical as the ratio between the degradation modes is constant. Path 1 (blue) had some exponential component for LLI and LAMPE and, therefore, the path is more complicated. Nonetheless, based on the triplets and the diagnosis dataset, the capacity loss, Figure 5(c), and the voltage variations, Figure 5(d), can be deciphered. This approach for calculating a prognosis dataset will be faster than simulating each degradation path one by one as it will avoid repeating the same simulations over and over.
Figure 5: Example of the prognosis dataset calculation for two different paths for a Gr/LFP cell with (a) the triplet variations, (b) their path in the ternary design space, (c) the reconstructed capacity loss, and (d) the voltage variations for path 1.

It must be noted that the description above was simplified for narrative purposes and was centered on the thermodynamic degradation observable at low currents and constant temperature for ease of representation. The full mechanistic approach is much more complex [25]. To be able to simulate different levels of usage, an equivalent circuit model (ECM) is included to handle different rates for each electrode [22-26]. For electrode blends, or changes in electroactive phases upon aging, each component of the electrode must be considered separately (i.e., LAMPE will be split in LAMLCO and LAMNCA for an LCO-NCA blended electrode). Moreover, for datasets at higher currents or different temperatures, some kinetic parameters such as the ohmic and faradic resistance increase must also be scanned as they will influence the voltage response of the cells. Finally, in some cases, lithium plating, both thermodynamic and kinetic, must be taken into consideration [25]. The validity of the mechanistic approach in all of those cases has been proven in the literature [19,58,59]. As explained in [25,58], resistance changes can be accommodated by shifting the voltage curves up or down depending on the electrode and the regime. Kinetic limitations can be handled by modifying the simulated rate while counterbalancing the polarization changes. Simulating higher rates allows emulating slower kinetics. Simulating lower rates allows emulating faster kinetics. This allows to model different rates and temperatures [58]. Figure 6(a) presents an example from [58] where different temperatures were simulated from 25°C data at different rates. For lithium plating, the solution is to simulate the NE as a blend between Gr and metallic lithium with the metallic lithium initially outside of the potential window, Figure 6(b). If LAMGraphite becomes prominent, or if its resistance pushes its potential below 0V, the metallic lithium will then be dragged into the potential window and start interacting with the PE [19,21,25]. As discussed in the literature [5,19,25,57], thermodynamic lithium plating begins when there is enough LAMNE so that the NE is fully lithiated before the PE is delithiated. The amount of LAMNE necessary for that to happen depends on the initial cell configuration (LRin and OFSin) and also on the evolution of LLI and LAMPE [60]. Figure 6(c) plots an example of the evolution of the plating threshold (PT) for a randomly chosen...
degradation path. Lithium plating will start when $\text{LAM}_{\text{NE}}$ becomes larger than $\text{PT}$. This does not by itself induce any capacity loss as metallic lithium can be reduced and oxidized. This lithium, however, is usually quickly passivating, which results in additional LLI. To take this effect into account, a correction for LLI can be introduced and the additional LLI is defined as the difference between $\text{LAM}_{\text{NE}}$ and the $\text{PT}$, when positive, multiplied by a reversibility factor. Figure 6(c) showcases an example of the acceleration in LLI based on different reversibilities. Figure 6(d) presents the associated capacity loss showcasing the acceleration.

![Image](image_url)

Figure 6: (a) Emulation of the impact of temperature at C/5 for a Gr//NCA cell from [58], and (b) Electrode matching with a blended Gr-Li electrode with the Li outside of the potential window as a reserve. Evolution of (c) LLI and $\text{LAM}_{\text{NE}}$ depending on the reversibility of lithium plating (0%, 50% or 100%) and (d) the associated capacity loss.

Proof-of-concept datasets

For this proof-of-concept work, a Gr//LFP cell similar to the one used in [20] was chosen because the sensibility analysis of different FOI was already reported and some non-ML diagnosis tools were validated. The data could also be compared to the one from Severson et al. [5].

The diagnosis dataset contains more than 500,000 voltage vs. capacity curves. It can be used to train machine learning algorithms, but more importantly, it can also be used to perform sensibility analysis [20] to find out which FOIs are representative of different degradations or if they can be used as universal proxies for capacity loss. An example is presented in Supplementary note 1.

The prognosis dataset contains more than 130,000 different degradation paths with upwards of 3,000,000 individual voltage vs. capacity curves. It can be used to test the validity of varying FOIs proposed in the literature for early prognosis. An example is presented in Supplementary note 2.

Both datasets are available to download (see Data availability section for access information).

Limitations and outlook

In this work, we presented a proof-of-concept methodology to generate big data training datasets for intercalation batteries. The mechanistic approach combines both modeling and experimental techniques to provide a universal tool for the creation of synthetic voltage curves practically indistinguishable from real data. This approach offers the benefits of the broad applicability of the model to various cell chemistries, designs, and operating modes, as well as the high fidelity as inhered from the detailed extraction of the experimental data. The methodology
could be applied to the generation of training datasets encompassing the entire degradation spectrum as well as different operating conditions such as rate and temperature. Scenarios leading to lithium plating with adjustable reversibility can be added and the addition of random noise on the data could be considered to make the datasets even more realistic and consider the impact of real-life testing machines’ precision and accuracy.

As a proof-of-concept, we generated a diagnosis dataset containing more than individual 500,000 voltage vs. capacity curves and a prognosis dataset with more than 130,000 individual degradation paths for a commercial graphite/LFP battery. These datasets are more than three orders of magnitude larger than the one currently available in the literature and they could be extended at will. We showcased the use of the proposed datasets for FOI sensibility analysis and a diagnosis technique based on the joint evolution of three FOIs since variations of single FOI were proven inefficient. We also used them to propose an early prognosis method based on the extrapolation of the diagnosis of the degradation modes.

Although there is still a lot of work to be done to optimize the technique, results are extremely promising and should accelerate the development of accurate algorithms. The proof-of-concept dataset used in this work were limited to thermodynamic degradations at constant temperature for single-phase electrodes. Taking everything into account (LAMs, LLI, polarization, kinetics, phases, temperatures…) for all the possible evolutions (linear, exponential, delays…) will take enormous calculation time and generate overwhelming datasets. Some optimization and intelligence could be applied to limit the number of degradation paths to test. A solution could be large designs of experiments first to screen the impact of the different parameters then to focus on their representativity in the paths to simulate. Here we provide an open framework to the community to apply advanced optimization techniques to reduce the calculation burden.

Finally, we would like to point out that our intent is no way to remove the need for experimental testing all-together. On the contrary, experimental testing is still essential, and the only way, to decipher which conditions cater to specific degradation paths and its input is, therefore, fundamental to decipher the applicability of batteries for a given application. Our methodology can be used alongside to develop the needed algorithms once a cell was selected.

Methods

Half-cell data. Data from four different types of commercial cells were used in this work. The Gr//NCA cell was a Panasonic 3350 mAh NCR 18650B [51,58,60-62]. The Si-Gr//NMC811 was a Samsung-SDI 3350 mAh INR18650-35E [52], the Gr//LCO+NCA was an LG-Chem 2800 mAh ICR18650 C2” [53], and the Gr//LFP was an A123 2300 mAh ANR26650M [19,63]. All were extensively studied in previous work. Interested readers are referred to the original publications for more details. The cells were slowly discharged to 0% SOC before being carefully disassembled in an Argon-filled glove box where the electrodes, separator, and casing will be separated [64]. Electrode discs (18 mm in diameter) were punched from the harvested electrodes and rinsed in a dimethyl carbonate (DMC) solution. The backside of two-sided electrodes was then wiped clean using cotton swabs soaked in n-N-methyl-2-pyrrolidone (NMP) solvent before being the electrode was rinsed again in a fresh DMC solution. The electrodes were tested using a lithium counter electrode. Metallic lithium was applied onto a stainless-steel disk. An electrolyte solution of 1.0 M LiPF6 in ethylene carbonate (EC) + DMC (1:1 by weight) + 2% wt. vinylene carbonate (VC) was used to soak the separator, which consisted of one layer of Whatman GF-D fiberglass discs (12.7 mm in diameter, Whatman, Kent, United Kingdom). Once sealed, the half-cells were taken out of the glove box and connected to a multi-channel Bio-Logic VMP3 potentiostat (Bio-Logic, Clai, France) for testing at different rates from low to high. The complete testing protocol can be found in [64].

Simulations. All simulations were performed under a MATLAB© environment using the ‘alawa toolbox [65]. As in [20], the LFP and graphite electrodes were matched with an LR of 0.95, OFS of 12.5%, and a resistance adjustment of -0.07 to match the A123 ANR26650M experimental data.
The diagnosis training dataset was compiled with a resolution of 0.01 for the triplets and C/25 charges. This accounts for more than 5,000 different paths at the base of the triangle in Figure 4. Each path was simulated with 0.85% increases for each degradation up to 85%. This accounts for 100 simulations per path. The training dataset, therefore, contains more than 500,000 voltage vs. capacity curves and took around 12h to compute on a standard laptop. The 500MB dataset is available to download (see Data availability section for access information).

The prognosis dataset was harder to define as there are no limits on how the three degradation modes can evolve. For this proof of concept work, we considered eight parameters to scan. For each degradation mode, degradation was chosen to follow equation (1).

\[
\%\text{degradation} = a \times \text{cycle} + (\exp^{b \times \text{cycle}} - 1) \quad (1)
\]

Considering the three degradation modes, this accounts for six parameters to scan. In addition, two other parameters were added, a delay for the exponential factor for LLI, and a parameter for the reversibility of lithium plating. The delay was introduced to reflect degradation paths where plating cannot be explained by an increase of LAMs or resistance [55]. The chosen parameters and their values are summarized in Table S1. Figure S1(a,b) presents the evolution of parameters \(p_1\) to \(p_7\). At the worst, the cells endured 100% of one of the degradation modes in around 1,500 cycles. Minimal LLI was chosen to be 20% after 3,000 cycles. This is to guarantee at least 20% capacity loss for all the simulations. For the LAMs, conditions were less restrictive, and, after 3,000 cycles, the lowest degradation is of 3%. The reversibility factor \(p_8\) was calculated with equation (2) when \(\text{LAM}_{\text{NE}} > \text{PT}\).

\[
\%\text{LLI} = \%\text{LLI} + p_8(\%\text{LAM}_{\text{PE}} - \text{PT}) \quad (2)
\]

Where \(\text{PT}\) was calculated with equation (3) from [60].

\[
\text{PT} = 100 - \left(\frac{100-\%\text{LAM}_{\text{PE}}}{100-\%\text{LAM}_{\text{PE}} - \%\text{LAM}_{\text{PE}}}\right) \times (100 - \%\text{OF}C_{\text{ini}} - \%\text{LLI}) \quad (3)
\]

Varying all those parameters accounted for more than 130,000 individual duty cycles. With one voltage curve for every 100 cycles, it took around 12h to compute on a standard laptop to simulate the more than 3,000,000 voltage vs. capacity curves. The 3GB dataset is available to download (see Data availability section for access information).

Data availability. The datasets used in this study are available at http://dx.doi.org/10.17632/bs2j56pn7y.1 and http://dx.doi.org/10.17632/6s6ph9n8zg.1 for the diagnosis and prognosis dataset, respectively.

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### Supplementary Tables and Figures

Table S1: Summary of scanned parameters for prognosis dataset

| Parameter | Description      | Values (% per cycle) |
|-----------|------------------|-----------------------|
| p1        | Linear coeff. LLI| 0.007, 0.010, 0.013, 0.017, 0.021, 0.027, 0.034, 0.048, 0.06 |
| p2        | Exp. Coeff. LLI  | 0.000001, 0.002, 0.0033 |
| p3        | Delay exp. LLI   | 500, 1000, 1500       |
| p4        | Linear coeff. LAMPE| 0.001, 0.005, 0.01, 0.015, 0.02, 0.025, 0.03, 0.0375, 0.05, 0.07 |
| p5        | Exp. Coeff. LAMPE| 0.000001, 0.001, 0.0013 |
| p6        | Linear coeff. LAMNE| 0.001, 0.005, 0.01, 0.015, 0.02, 0.025, 0.03, 0.0375, 0.05, 0.07 |
| p7        | Exp. Coeff. LAMNE| 0.000001, 0.001, 0.0013 |
| p8        | Plating reversibility| 0, 50, 100           |

Figure S1: Different duty cycles tested for each degradation mode with (a) LLI and (b) the LAMs.
Supplementary Note 1: Diagnosis Dataset Analysis

Figure S2 presents an example of the sensibility of 3 FOIs for capacity loss, LLI, LAMfr, and LAMne. The three selected FOIs were the capacity of the first plateau (area under peak ① on Figure S3), the capacity of everything but the first plateau (area under peaks ②-⑤), and the capacity of the last plateau (area under peak ⑤) [20]. As reported in our previous work, [20], the sensibility analysis showcase that no single FOI can track capacity loss nor one of the degradation modes alone since there is always a spread of values. Nonetheless, as reported in [20], their combined variations might be used. This will be discussed in supplementary note 2.

Figure S2: Sensibility analysis for 3 FOI on the Gr/LFP C/25 charges.

Figure S3: IC signature of a Gr/LFP C/25 charge with the typical peak indexed 1-5.
Supplementary Note 2: Prognosis Dataset Analysis

The prognosis dataset can be used to test the validity of different FOIs proposed in the literature for early prognosis. Figure S4 (a,b) presents the evolution of the variance between cycle one and cycle 100 and 500 respectively as a function of cycle life, defined as the cycle at which 20% capacity loss is reached. When considering a multitude of degradation paths, the variance approach seems less effective than proposed in the literature [5,11] with correlation coefficients of -0.56 and -0.43, respectively, when calculated after 100 and 500 cycles. Looking at the capacity loss after 100 cycles, Figure S4 (c), the correlation is better (ρ = 0.63). Still, the error could be huge for low capacity loss as paths with less than 1% capacity loss after 100 cycles were found to induce 20% capacity loss after just 200 cycles. Finally, as predicted [20,21], the evolution of the first LFP plateau capacity (i.e., the area under peak ① in the Gr//LFP IC curves) is not a good indicator with a Pearson coefficient of 0.51, Figure S4(d). The dataset could be analyzed further by tracking if specific conditions exist where the FOI is accurate, e.g., the first LFP plateau capacity is likely working well when there is little LAMNe [20,21]. This is out of the scope of this publication and will be investigated in future work.

The prognosis dataset can also be used to validate diagnosis techniques. For example, the method proposed in [20] can be tested. As shown in Figure S2 individual FOI information is not enough to reach a proper diagnosis, but their combined variations might be enough. Figure S5 presents the diagnosis results based on the combined variations of 3 FOIs for diagnosis after 100, 200, 500, and 1,000 cycles for the 130,000 computed duty cycles. Out of the more than 500,000 points tested (4 x 130,000), the average errors for estimation of LLI, LAMRe, LAMNe, and capacity loss were of -0.32%, -2.73%, -0.46%, and -0.33% respectively with standard deviations below 1% for all but LAMRe (6%) for diagnosis up to the 500th cycle. The distribution of the results is presented in Figure S5 and the different statistics summarized in Table S2. For all, but LAMRe, most of the errors are comprised between -1 and 1%, with the minimum average error recorded for the diagnosis after 500 cycles and the maximum for the diagnosis after 1,000 cycles. The spread for LAMRe is larger, between -2 and 10%, which was expected since LAMRe is notoriously hard to quantify for LFP cells because of the single voltage plateau [25]. The average LAMRe error decreased with cycle number to 0.74% after 1,000 cycles. Looking more into details, the maximum and minimum errors were always recorded for the diagnosis at 1,000 cycles with maximum and minimum errors mostly below 10% for all the other cycles (except for LAMRe). Future work will investigate the specific combinations that led to large errors, although our previous work [20] showed that it was for degradations unlikely to happen under real-life conditions where LLI usually predominates.
Figure S4: Evolution of the (a) the variance between cycle 100 and 1, (b) the variance between cycle 500 and 1, (c) the capacity loss at cycle 100 and (d) the area of the high voltage IC peak at cycle 100 as a function of cycle life (i.e., cycle at which 20% capacity loss is reached).

Figure S5 Distribution of the estimation errors for (a) LLI, (b) LAMPE, (c) LAMPE, and (d) the capacity loss based on the evolution of 3 FOIs at cycle 100, 200, 400, and 1000.
Table S2: Main error with standard deviation as well as minimum and maximum error for the three degradation modes and the capacity loss.

|        | Mean (%) | Std (%) | Minimum (%) | Maximum (%) |
|--------|----------|---------|-------------|-------------|
| LLI    | -0.32    | 2.39    | -71.86      | 39.31       |
| cycle 100 | -0.12    | 0.34    | -1.53       | 0.87        |
| cycle 200 | -0.08    | 0.36    | -1.42       | 1.45        |
| cycle 500 | -0.02    | 0.79    | -8.73       | 9.39        |
| cycle 1000 | -1.13    | 4.75    | -71.86      | 39.31       |
| LAM$\text{PE}$ | 2.73     | 5.82    | -61.35      | 63.60       |
| cycle 100 | 4.17     | 2.71    | -3.66       | 13.07       |
| cycle 200 | 3.47     | 3.47    | -8.82       | 21.05       |
| cycle 500 | 2.31     | 5.84    | -34.10      | 41.15       |
| cycle 1000 | 0.74     | 8.98    | -61.35      | 63.60       |
| LAM$\text{NE}$ | -0.46    | 2.36    | -49.91      | 46.52       |
| cycle 100 | -0.45    | 0.42    | -1.65       | 0.85        |
| cycle 200 | -0.39    | 0.40    | -1.97       | 1.00        |
| cycle 500 | -0.28    | 0.67    | -11.49      | 4.28        |
| cycle 1000 | -0.75    | 4.78    | -49.91      | 46.52       |
| Capacity loss | -0.33    | 2.37    | -60.43      | 43.29       |
| cycle 100 | -0.18    | 0.35    | -1.32       | 1.16        |
| cycle 200 | -0.10    | 0.41    | -1.51       | 1.86        |
| cycle 500 | -0.03    | 0.92    | -10.20      | 10.68       |
| cycle 1000 | -1.04    | 4.69    | -60.43      | 43.29       |

The results from Figure S4 raise the important question of whether a single point early prognosis is possible. With the diagnosis approach validated, a possible solution could be to use different diagnoses and extrapolate the evolution of the different parameters to reconstruct the voltage curves using the mechanistic approach, Figure S6. This technique is already used to forecast the evolution of capacity loss for large experimental studies [59,61,66]. Further work is in progress to affine the technique further and to understand under which conditions the estimation is the most accurate.

Figure S6: (a) Schematic representation of possible early prognosis solution using the mechanistic approach.