# Alternative Approach to the Holomorphic Embedding Load-Flow Method

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Abstract—The Holomorphic Embedding Load-Flow Method is a novel recursive technique for solving the power flow of an electrical power system. Contrary to the traditional iterative schemes, it embeds the equations in a bigger problem from where the solution is then built. There is a wide range of possibilities to adapt the equations to the holomorphic embedding method. Although the so-called canonical embedding prevails as the most popular one, achieving the desired reference state may result in overcomplicating the algorithm. This paper covers a simple alternative embedding, capable of offering adequate convergence properties, sometimes even faster than the canonical embedding. An 11-bus ill-conditioned system is studied in order to show that improvement. Furthermore, despite the fact that unique tools such as Sigma and Thévenin approximants were constructed with the canonical embedding in mind, they become fully compatible with the alternative embedding thanks to minor modifications. They are applied to the IEEE 30-bus system to display its functionality.

*Index Terms*—holomorphic embedding method, load flow, power system analysis computing, power system simulation, Sigma approximants, Thévenin approximants.

# I. INTRODUCTION

POWER flow or load flow is likely the most relevant problem concerning power systems. It describes the steadystate operation of the grid, which means that voltages, currents and powers are completely known. Unfortunately, it is a wellknown fact that the equations involved are nonlinear. Because of that, power flow calculations have been usually performed by iterative methods, such as the Gauss-Seidel and mostly the Newton-Raphson and its variations [1], [2], with the fast decoupled load flow being one of them [3]. The Newton-Raphson method converges quadratically. That causes it to converge to an acceptable solution with few iterations.

However, iterative methods are not guaranteed to always obtain the right solution. There are two main problems that the Newton-Raphson method and similar approaches have to face: one being the divergence of the solution, and the other, the convergence to an infeasible operating point [1].

That is where the holomorphic embedding method comes into play. When the power flow is solvable, the method reaches the correct solution, whereas when the power flow is impossible to solve, the method signals it [4], [5], [6]. The holomorphic embedding is based on, as the name suggests, submerging the original problem in a larger problem. When

some reference conditions are provided, it is then able to compute the solution [5].

The choice of embedding is not unique [1], [5], [7], [8]. The most notorious one looks for a reference state where the first coefficients of the unknowns are always the same, independent of the system [5], [9], [10]. It will be referred to as the canonical embedding. Despite its relative complexity, not only it works properly, but it is also compatible with the totality of tools that the holomorphic embedding method has to offer. Probably the most prominent tools are Sigma and Thévenin approximants, detailed in [5].

Some authors have opted for an approach slightly less complex [1], [8], [11]. The main trouble has to do with PV buses, where the injected reactive power is unknown. References [8], [12] worked with an implementation that manages to only consider the voltages as unknowns. Nonetheless, that embedding lacks robustness, as it presents convergence problems. That has been affirmed in [1], where the initialization is carried out with iterative methods, but in essence the procedure stays the same.

To solve that, several embeddings are developed in [13]. They successfully achieve smaller errors than [11], [12]. Despite that, it can be argued that the embedding is somewhat complex, or at least, way different from the canonical embedding. The effects of tap changers are not mentioned and neither unique tools of the holomorphic embedding method like Sigma and Thévenin approximants are covered.

This paper details an approach similar to the canonical embedding which turns out to be slightly simpler. Contrary to [1], [8], [11] its convergence properties are satisfactory, and in certain situations even better than the canonical embedding. Another advantage is its compatibility with Sigma and Thévenin approximants.

The outline of this work is as follows: Section II introduces the canonical embedding; Section III presents the alternative embedding and details the algorithm that has to be employed; Section IV reviews the concept and the importance of Sigma approximants, describes the modification needed in order to make it compatible with the alternative embedding and then, compares it with the results obtained with the canonical embedding; Section V depicts the formulation of Thévenin approximants, covers the changes and the steps that need to be taken as well as showing its usage and how well they match the values resulting from the canonical embedding; finally, Section VI concludes the paper.

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## II. CANONICAL EMBEDDING

The Holomorphic Embedding Load-Flow Method takes an initial equation of the form

$$\sum_{j} Y_{ij} V_j + Y_i^{sh} V_i = \frac{S_i^*}{V_i^*},\tag{1}$$

where  $Y_{ij}$  is the i,j element of the series admittances bus matrix, V stands for voltage,  $Y_i^{sh}$  represents the object that collects all shunt admittances and  $S_i$  is the power injected at bus i. Current injections could also be considered, although they have been ignored with the idea to keep the algorithm simple.

The idea behind the embedding is based on integrating the power flow problem into a much larger problem. Under these circumstances, the construction of the solution initiates from a well-defined reference state, from which one is then able to extend the solutions [5]. The traditional embedding transforms (1) into

$$\sum_{j} Y_{ij} V_{j}(s) + s Y_{i}^{sh} V_{i}(s) = s \frac{S_{i}^{*}}{V_{i}^{*}(s^{*})}, \tag{2}$$

where s is the complex variable and  $V_i(s)$  is in essence a series that obeys the generic form

$$C(s) = C[0] + sC[1] + s^2C[2] + \dots + C[n] = \sum_{k=0}^{n} s^kC[k], (3)$$

where n represents an arbitrary depth, usually between 20 and 40. One has to calculate all terms step by step. Not only voltages, but also each and every unknown variable follows the structure of (3). The holomorphic embedding starts from a reference state, where s=0, and then, it looks for the final solution at s=1.

While it is true that the embedding can be chosen freely, it has to validate the Cauchy-Riemann conditions. Due to that specific reason, the last term of (2) is embedded as  $V_i^*(s^*)$  and not as  $V_i^*(s)$ . Both the canonical and the alternative embedding described in this paper adhere to that.

At this point not only PQ buses should be considered. In addition to them, power systems contain PV buses, where reactive power is treated as an unknown. The original embedded equations follow

$$\sum_{j} Y_{ij} V_j(s) + s Y_i^{sh} V_i(s) = \frac{s P_i - j Q_i(s)}{V_i^*(s^*)}, \tag{4}$$

where  $P_i$  is the given active power and  $Q_i(s)$  becomes the series with the reactive power coefficients. It has to be noted that the term  $Q_i(s)$  does not multiply by s. That is critical because the initial solution (the one corresponding to the reference state) should emerge from a linear system of equations.

The way to deviate from the mentioned inconvenience consists in establishing beforehand

$$\begin{cases} V_i[0] = 1 & i \in PQ, PV, \text{ slack}, \\ Q_i[0] = 0 & i \in PV, \end{cases}$$
 (5)

where PQ, PV and slack are the sets containing their respective type of bus.

With the aim of forcing the initialization of  $V_i[0]=1$  it is mandatory to work with an admittance bus matrix where the sum of all elements of every single row becomes 0. However, that does not hold true for transformers working with offnominal tap ratios. The admittance bus matrix could remain symmetrical given that no phase changes are introduced. In spite of that, as soon as the transformation ratio differs from the unit, rows are not guaranteed to sum exactly 0.

One way to confront the problem is based on dividing the admittance bus matrix in two: one that mimics the full matrix but establishes that transformation ratios are equal to  $1(Y^{(b)})$  and a matrix that is meant to cover the influence of tap changers  $(Y^{(a)})$ . They are combined into

$$Y = sY^{(a)} + Y^{(b)}. (6)$$

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The sum of both matrices is equal to the original admittance bus matrix Y. Multiplying the matrix  $Y^a$  by s manages to neglect its effects on the reference state so that only the  $Y^{(b)}$  matrix has to be taken into account at first. That way, (5) becomes validated. Reference [14] exemplifies this division of matrices in a given two-bus link.

There is one last expression necessary with PV buses

$$V_i(s)V_i^*(s^*) = 1 + s(W_i - 1), \tag{7}$$

where  $W_i$  is the squared absolute value of the voltage of bus i. That expression is only valid when the first voltage coefficients are 1.

## III. ALTERNATIVE EMBEDDING

In contrast to the canonical embedding, the alternative embedding is not concerned with forcing that all voltage coefficients are 1 as well as having all reactive power coefficients equal to 0 in the beginning. In fact, it is perfectly fine to initiate the series with voltages slightly different to 1 and at the same time different from one and the other.

The embedding of PQ buses does not differ from (2). However, the novelty comes with the PV buses equations. On the one hand, (4) transforms into

$$\sum_{i} Y_{ij} V_j(s) + s Y_i^{sh} V_i(s) = s \frac{P_i - j Q_i(s)}{V_i^*(s^*)}.$$
 (8)

The only difference is that the reactive power is also multiplied by s, which makes the whole term on the right-hand side of the equation unimportant to the reference state. Thus, voltage coefficients of PQ and PV buses are initialized by solving directly a linear system of equations.

With the presence of tap changers, the first voltage coefficients are not exactly 1, but usually, similar to it. More than an inconvenience, that becomes a benefit. Voltages coefficients can be closer to the final solution just from the very first order, which can positively affect the convergence rate. Moreover, the bus admittance matrix does not have to be split into two parts, which simplifies the algorithm.

The embedding selected for the slack bus voltage  $(V_w(s))$  may be identical to the one used in the canonical embedding

$$V_w(s) = 1 + s(V_w - 1), (9)$$

where  $V_w$  is a given value. Despite that, it is also functional to not define it this way.

When it comes to building the solution of the reactive power series, their firsts terms are obtained at the same time that the second coefficients of the voltage series are calculated, and so on. Their calculation is always one step behind, although that carries no tragic consequences.

Finally, (7) of the canonical embedding becomes

$$V_i(s)V_i^*(s^*) = V_i[0]V_i^*[0] + s(W_i - V_i[0]V_i^*[0]).$$
 (10)

In the canonical embedding, that equation did not participate in defining the reference state, but it had to be consistent with it. Similarly with the alternative embedding, even though now the firsts coefficients may differ from 1. Hence the given change.

## A. Calculation of terms

As it has been announced, all the terms that form the unknowns have to be obtained. First, at the reference state, (10) is ignored and the first voltage coefficients appear from solving the following system of linear equations

$$\sum_{j \neq w} Y_{ij} V_j[0] = -Y_{iw}, \tag{11}$$

where  $Y_{iw}$  is the admittance bus matrix element that connects the i bus to the slack bus. Once the first voltage coefficients are computed, since the calculation of the reactive power terms is delayed, there are no variables left to find. Therefore, the reference state becomes completely defined.

For the next coefficients, it is convenient to introduce the change of variable

$$X_i(s) = \frac{1}{V_i^*(s^*)}. (12)$$

As a consequence of that, its coefficients are calculated as

$$X_{i}[c] = \begin{cases} \frac{1}{V_{i}^{*}[c]} & c = 0, \\ \frac{-\sum_{k=0}^{c-1} X_{i}[k]V_{i}^{*}[c-k]}{V_{i}^{*}[0]} & c \ge 1, \end{cases}$$
(13)

which only depends on the voltage coefficients of the same or inferior order. In that regard, this is independent of the chosen embedding.

Now, the second voltage coefficients and the first reactive power coefficients have to be found. The reader may refer to [15] to discover how the expansion of coefficients works in more detail (even though it covers the canonical embedding). For PQ buses (2) becomes

$$\sum_{j\neq w} \left( G_{ij} V_j^{(re)}[1] - B_{ij} V_j^{(im)}[1] \right) =$$

$$\Re[Y_{iw}(V_w - 1) - Y_i^{sh} V_i[0] + S_i^* X_i[0]],$$

$$\sum_{j\neq w} \left( B_{ij} V_j^{(re)}[1] + G_{ij} V_j^{(im)}[1] \right) =$$

$$\Im[Y_{iw}(V_w - 1) - Y_i^{sh} V_i[0] + S_i^* X_i[0]],$$
(14)

where  $G_{ij}$  and  $B_{ij}$  are respectively the real and imaginary part of the admittance element  $Y_{ij}$ . References [1], [8], [11] do not split the expression between real and imaginary components.

The algorithm is considerably less complex, however, it has been stated that convergence properties may suffer from that.

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As for PV buses, (8) originates

$$\sum_{j\neq w} \left( G_{ij} V_j^{(re)}[1] - B_{ij} V_j^{(im)}[1] \right) - X_i^{(im)}[0] Q_i[0] =$$

$$\Re[Y_{iw}(V_w - 1) - Y_i^{sh} V_i[0] + P_i X_i[0]],$$

$$\sum_{j\neq w} \left( B_{ij} V_j^{(re)}[1] + G_{ij} V_j^{(im)}[1] \right) + X_i^{(re)}[0] Q_i[0] =$$

$$\Im[Y_{iw}(V_w - 1) - Y_i^{sh} V_i[0] + P_i X_i[0]],$$
(15)

where the first reactive power coefficients become involved. Voltages absolute values, which are defined by (10), in terms of coefficients turn into

$$2V_i^{(re)}[0]V_i^{(re)}[1] + 2V_i^{(im)}[0]V_i^{(im)}[1] = W_i - |V_i[0]|^2.$$
 (16)

As a result of that, a linear system of equations (built by (14), (15) and (16)) has been derived. In compact form the system becomes

$$\begin{pmatrix} G & B & -X^{(im)}[0] \\ B & G & X^{(re)}[0] \\ 2V^{(re)}[0] & 2V^{(im)}[0] & 0 \end{pmatrix} \begin{pmatrix} V^{(re)}[1] \\ V^{(im)}[1] \\ Q[0] \end{pmatrix} = \begin{pmatrix} RHS(14) \\ RHS(15) \\ RHS(16) \end{pmatrix},$$
(17)

where RHS stands for the right-hand side expression of (14), (15) and (16). This matrix structure becomes similar to the model 3 of [13], although the imaginary part of the voltages may not be null and the PV voltage absolute value differs. Just like the canonical embedding, the matrix involved remains constant and can also be considered sparse. In the alternative embedding, it tends to be slightly denser, albeit these characteristics contribute to reducing the computational time.

From now on the algorithm can be generalized. It is relevant to note that the canonical embedding would require to derive the equations for another step before being extended for any given order of coefficients. Considering  $c \geq 2$ , which represents the current step, (2) is converted to

$$\sum_{j \neq w} \left( G_{ij} V_j^{(re)}[c] - B_{ij} V_j^{(im)}[c] \right) = \Re[-Y_i^{sh} V_i[c-1] + S_i^* X_i[c-1]],$$

$$\sum_{j \neq w} \left( B_{ij} V_j^{(re)}[c] + G_{ij} V_j^{(im)}[c] \right) = \Im[-Y_i^{sh} V_i[c-1] + S_i^* X_i[c-1]].$$
(18)

Voltage absolute values of PV buses obey

$$2V_i^{(re)}[0]V_i^{(re)}[c] + 2V_i^{(im)}[0]V_i^{(im)}[c] = -\sum_{k=1}^{c-1} V_i[k]V_i^*[c-k].$$
(19)

Additionally, (8) gives place to

$$\sum_{j\neq w} \left( G_{ij} V_j^{(re)}[c] - B_{ij} V_j^{(im)}[c] \right) - X_i^{(im)}[0] Q_i[c-1] = 
\Re\left[ -j \sum_{k=1}^{c-1} X_i[k] Q_i[c-1-k] - Y_i^{sh} V_i[c-1] \right] 
+ P_i X_i[c-1],$$

$$\sum_{j\neq w} \left( B_{ij} V_j^{(re)}[c] + G_{ij} V_j^{(im)}[c] \right) + X_i^{(re)}[0] Q_i[c-1] = 
\Im\left[ -j \sum_{k=1}^{c-1} X_i[k] Q_i[c-1-k] - Y_i^{sh} V_i[c-1] \right] 
+ P_i X_i[c-1].$$
(20)

Thus, the algorithm has been fully generalized. In order to calculate the following coefficients it is enough to use (18), (19) and (20) in combination with the matrix of (17) to build several linear systems of equations. Building and solving them sequentially generates the terms. As a matter of fact, the procedure is basically the same that one encounters with the canonical embedding.

The next step to find out the solutions consists in evaluating the series at s=1. Several possibilities could be contemplated to achieve so [5]. Padé approximants are usually the go-to choice because of Stahl's theorem, even though recursive methods such as Wynn's  $\varepsilon$  or Bauer's  $\eta$  [16], [17] are also suitable choices [9]. That last one resource is not totally appropriate for the canonical embedding, but offers no inconvenience with the alternative embedding due to having the first reactive power coefficients different from zero.

## B. Results

The alternative embedding has been tested for the next grids: IEEE 14, IEEE 30 and IEEE 118-bus system. In all of them, both approaches (the canonical and the alternative embedding) were capable of solving the grids at similar convergence rates. Table I displays the smallest radius of convergence of the series involved in every grid.

TABLE I
MINIMUM RADIUS OF CONVERGENCE WITH BOTH EMBEDDINGS

	Embedding		
Grid	Canonical	Alternative	
IEEE 14	3.68	4.53	
IEEE 30	5.82	5.72	
IEEE 118	2.64	1.94	

Yet, there was an extra grid in which the alternative embedding was superior to the canonical embedding. That corresponds to an ill-conditioned 11-bus system [18], [19]. The radius of convergence were 0.88 and 2.84 respectively. Out of 14 branches, 7 contain transformer tap changers. A loading factor of 0.5 has been chosen in order to unload the system. Despite that, the basic Newton-Raphson method converged to

low-voltage solutions, which from an operation standpoint, are incorrect.

Under these conditions, Fig. 1 plots the logarithm of the maximum complex power error versus the number of coefficients picked. The final results were generated with the near-diagonal Padé approximants.

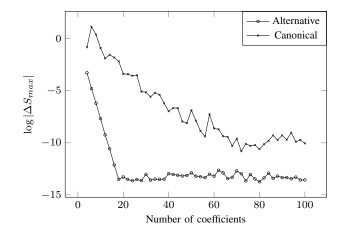


Fig. 1. Maximum errors depending on the number of coefficients for the 11-bus system.

In the beginning, the error obtained by the alternative embedding decays linearly in the logarithmic scale. Then, it reaches a plateau, which is to be expected [5]. It draws a shape that resembles a hockey stick. Double-precision was used. On the other hand, the alternative embedding progresses much more slowly. From the start the errors are bigger, and while they tend to be reduced when the number of coefficients increases, around 70 terms are needed to reach an error of about  $10^{-10}$ . From then on, it does not improve.

Of course, in that case the solution could be refined thanks to the Padé-Weierstrass procedure [5] so that it would eventually reach the maximum attainable precision and compete with the originally given by the alternative embedding. Nonetheless, in that particular case, the alternative embedding on the start is more suitable. It does not force V[0] = 1, so that the first coefficients can approach with much greater accuracy the final value. For instance, bus number 11 presents a final voltage of 1.759; the first coefficient turns out to be 1.650.

## IV. INTEGRATION OF SIGMA APPROXIMANTS

Sigma approximants were introduced in [6] as a diagnostic tool. That is a crucial advantage that the holomorphic embedding possesses over the iterative methods such as the Newton-Raphson. Sigma approximants are able to show if the solution is feasible or infeasible, and furthermore, tell qualitatively where the problem lies. They offer graphical results, from which the human eye can extract patterns.

## A. Formulation

The definition of Sigma approximants starts from considering a two-bus equivalent between a given bus i and the slack bus

$$\frac{V_i(s) - V_w(s)}{Z_{iw}(s)} = s \frac{S_i^*(s^*)}{V_i^*(s^*)},\tag{21}$$

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where  $Z_{iw}(s)$  can be thought of equivalent to an impedance between both buses and  $S_i(s)$  represents the power injection at bus i. The procedure that follows is not concerned to find them individually, but rather obtain  $\sigma(s)$ , which will combine  $Z_{iw}(s)$  and  $S_i^*(s^*)$ .

The development of (21) starts by multiplying on both sides of the expression by  $Z_{iw}(s)/V_w(s)$ . As a consequence

$$\frac{V_i(s)}{V_w(s)} - 1 = s \frac{S_i^*(s^*) Z_{iw}(s)}{V_i^*(s^*) V_w(s)}.$$
 (22)

For commodity purposes, a new variable is defined

$$U_i(s) = \frac{V_i(s)}{V_w(s)},\tag{23}$$

which illustrates the ratio between the chosen bus of study and the slack bus. Combining (23) and (22) yields

$$U_i(s) - 1 = s \frac{S_i^*(s^*) Z_{iw}(s)}{V_i^*(s^*) V_w(s)} \frac{V_w^*(s^*)}{V_w^*(s^*)}.$$
 (24)

From here the definition of  $\sigma(s)$  becomes

$$\sigma(s) = \frac{S_i^*(s^*)Z_{iw}(s)}{V_{w}(s)V_{s*}^*(s^*)}.$$
 (25)

As it has been mentioned, the slack bus voltage series is captured in (9) and is fully known. The series  $Z_{iw}(s)$  and  $S_i^*(s^*)$  are unknown. Despite that, the procedure to calculate  $\sigma(s)$  makes use of voltage coefficients and not of these series. Eventually,  $\sigma(s)$  participates in the two-bus equivalent

$$U_i(s) = 1 + s \frac{\sigma(s)}{U_i^*(s^*)}. (26)$$

An important consequence is derived from (26): by definition,  $U_i[0] = 1$ . However, that is not always the case with the alternative embedding. It is likely that as a consequence of the introduction of tap changers the first voltage coefficients are not exactly 1, and because the slack bus is embedded as (9),  $U_i[0] \neq 1$ . This invalidates the usage of Sigma approximants.

To solve that problem, it is enough to transform (26) into

$$U_i(s) = 1 + \frac{\sigma'(s)}{U_i^*(s^*)},$$
 (27)

where  $\sigma'(s)$  is the newly defined series. The reason that explains why this transformation works lies in the fact that  $\sigma(s)$  is evaluated at s=1, and precisely (26) as a whole also is. Therefore, one can expect that  $\sigma(s=1)=\sigma'(s=1)$ .

It is straightforward to express the ratio of voltages as a function of  $\sigma'^{(re)}$  as well as  $\sigma'^{(im)}$ , which symbolize the real and imaginary part of  $\sigma'(s=1)$ 

$$U = \frac{1}{2} \pm \sqrt{\frac{1}{4} + \sigma'^{(re)} - (\sigma'^{(im)})^2} + j\sigma'^{(im)}.$$
 (28)

Therefore, the power-flow solution is incorrect if the discriminant turns negative, just as it happens with the original definition of  $\sigma(s)$ .

In turn, it is favorable to calculate  $\sigma'(s)$  through a rational function. From (27)

$$\frac{U_i(s) - 1}{1/U_i^*(s^*)} = \sigma'(s). \tag{29}$$

In essence, this procedure is equivalent to the Padé approximants.  $\sigma'(s=1)$  comes from solving a linear system. An alternative to that is based on using osculation relations [5].

## B. Results

Fig. 2 shows the Sigma plot for the IEEE 30 bus system with an increase of active power load from 0.106 to 0.82 at bus number 30. The voltage series involved 60 coefficients and Padé approximants were obtained with the matrix method, detailed in [5], [9].

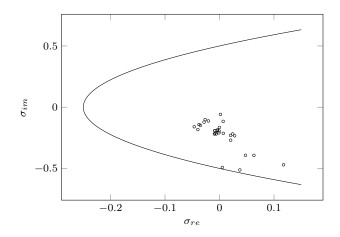


Fig. 2. Sigma plot of the IEEE 30 bus system under stressed conditions.

It is noticeable that a couple of buses approach the limit of the parabola. They are buses number 30 and 29. The first one of them suffers, in fact, the increase in load, whereas bus number 29 connects directly to it. By looking at the Sigma it is possible to identify what area of the system is under more stress.

The results obtained with the canonical embedding practically speaking do not differ from the ones provided by the alternative embedding. Table II represents the maximum absolute and relative difference between  $\sigma(s=1)$  and  $\sigma'(s=1)$  given several loads on bus number 30. The results have been generated working with double-precision floating-point format.

TABLE II  $\mbox{Maximum errors between } \sigma(s=1) \mbox{ and } \sigma'(s=1)$ 

-0.25     1.90E-15     1.78E-13       -0.50     9.07E-15     6.53E-14       -0.75     7.62E-12     6.14E-11       -0.82     6.03E-05     1.51E-04	P <sub>30</sub>	$ \sigma(s=1) - \sigma'(s=1) $	$\frac{ \sigma(s=1) - \sigma'(s=1) }{ \sigma(s=1) }$
-0.75 7.62E-12 6.14E-11	-0.25	1.90E-15	1.78E-13
****	-0.50	9.07E-15	6.53E-14
-0.82 6.03E-05 1.51E-04	-0.75	7.62E-12	6.14E-11
	-0.82	6.03E-05	1.51E-04

More than errors, these values should be understood as differences, since no evidence supports that  $\sigma(s=1)$  is a finer solution in comparison to  $\sigma'(s=1)$ . In any case, the differences are relatively small. It has to be taken into account that the voltage collapse point has been found around  $P_{30}=-0.8252$ . Thus, the values obtained for  $P_{30}=-0.82$  are extremely close to the voltage collapse point, so these differences are not surprising. Overall it can be stated that the

adaptation of Sigma approximants is appropriate and yields satisfactory results.

## V. INTEGRATION OF THÉVENIN APPROXIMANTS

Thévenin approximants consist in a particular case of Hermite-Padé approximants [5]. They approximate the voltage of a certain bus thanks to the osculation relation, which is defined as

$$T_N^{(0)}(s) + T_N^{(1)}(s)V(s) + T_N^{(2)}(s)\frac{s}{V^*(s^*)} = r_N(s)s^{N+1}, (30)$$

where expressions of the form  $T_N^{(a)}$  are polynomials to be found and  $r_N$  is the residue at a given order  $N=-1,0,1,...,\infty$ . Note that the subindex referring to a concrete bus is omitted.

The idea behind (30) is to define a resembling two-bus equivalent, hence the name of Thévenin approximants. Organizing (30) the voltage series becomes

$$V(s) = -\frac{T_N^{(0)}(s)}{T_N^{(1)}(s)} - \frac{T_N^{(2)}(s)}{T_N^{(1)}(s)} \frac{s}{V^*(s^*)} + r_N'(s)s^{N+1}, \quad (31)$$

which in some sense reminds of (27). Similar to the aforementioned Sigma approximants, it is useful to compact (31) as

$$V(s) = V^{(sw)}(s) + s \frac{S^{(Th)}(s)}{V^*(s^*)},$$
(32)

where  $V^{(sw)}(s)$  depends on the quotient of  $T_N^{(0)}(s)$  and  $T_N^{(1)}(s)$ . It can be understood as some kind of voltage tagged to the slack bus.  $S^{(Th)}$  illustrates the equivalent complex power injected into the given bus. It is calculated from the quotient of  $T_N^{(2)}(s)$  and  $T_N^{(1)}(s)$ .

of  $T_N^{(2)}(s)$  and  $T_N^{(1)}(s)$ .

Moreover, by adding the variables  $u=\frac{V}{V^{(sw)}}$  and  $\sigma^{(Th)}=\frac{S^{(Th)}}{V^{(sw)}(V^{(sw)})^*}$ , the solution of u is obtained by

$$u = \frac{1}{2} \pm \sqrt{\frac{1}{4} + \Re[\sigma^{(Th)}] - (\Im[\sigma^{(Th)}])^2} + j\Im[\sigma^{(Th)}], \quad (33)$$

which at its core is identical to (28). Sigma approximants where not used to compute the final voltage solution using (28) (although they could be employed for this exact purpose). In contrast, Thévenin approximants are used to find the final voltage solution. Once both solutions of u are obtained, the product of them by  $V^{(sw)}$ , which becomes a known series, yields the final values of V.

On the one hand, Thévenin approximants can be used to be certain about where the voltage collapse point lies. When the discriminant becomes negative, no solution is valid. On the other hand, they provide the user with the solutions of both stable and unstable operation. In other words, thanks to them one is able to construct PV and QV curves.

## A. Calculation process

The calculation of Thévenin approximants has been covered in [5]. However, the first voltage coefficients were assumed to be always 1. As a consequence of that, it is necessary to extend the calculation procedure independently of what the first coefficient turns out to be.

First of all, the condition concerning the degree of polynomials of the form  $T_N^{(a)}(s)$ , where a=0,1,2, is

$$\operatorname{degree}\left(T_N^{(a)}(s)\right) = \left|\frac{N+1-a}{3}\right|. \tag{34}$$

As a byproduct of that, when N increases by 1, only one of the three polynomials increments its degree. When the degree of a polynomial is negative, the polynomial becomes null.In case the degree is 0, the polynomial becomes a constant. These are precisely the two cases one encounters during the initialization.

The normalization condition that follows is an arbitrary choice. It is expected that the first voltage coefficients are not so distant from 1. Thereferore

$$T_N^{(0)}(0) = -1, (35)$$

where it is irrelevant what N it refers to. Note that this condition is identical to the chosen in [5], without loss of generality. As it can be deduced from (30) and (34), the next set of polynomials result in

$$\begin{cases}
T_{-1}^{(1)} = 0, \\
T_{0}^{(1)} = \frac{1}{V[0]}, \\
T_{1}^{(1)} = \frac{1}{V[0]}.
\end{cases}$$
(36)

Lastly, the next and final set of polynomials to initialize become

$$\begin{cases}
T_{-1}^{(2)} = 0, \\
T_{0}^{(2)} = 0, \\
T_{1}^{(2)} = \frac{-V[1]V^{*}[0]}{V[0]}.
\end{cases}$$
(37)

At this stage all polynomials could be understood as constants. However, that may not be the case in the next orders. Before calculating them, it is required to initialize the residues

$$\begin{cases}
r_{-1}[0] &= -1, \\
r_{0}[k] &= \frac{V[k+1]}{V[0]}, \\
r_{1}[k] &= \frac{V[k+2]}{V[0]} - \frac{V[1]V^{*}[0]X[k+1]}{V[0]},
\end{cases} (38)$$

where k=0,1,...,n-2, n represents the last index of the series V(s) and  $X(s)=\frac{1}{V^*(s^*)}$ .

The goal now is to increase N as much as possible, so the polynomials are able to approximate the two voltage solutions at its maximum precision. In order to progress towards the next step, which is N+1, new polynomials are defined as

$$T_{N+1}^{(a)}(s) = a_{N+1}sT_{N-2}^{(a)} + b_{N+1}T_{N-1}^{(a)} + c_{N+1}T_N^{(a)}, \quad (39)$$

where constants  $a_{N+1}$ ,  $b_{N+1}$  and  $c_{N+1}$  have to be found. From now on the procedure is exactly the same as [5]. The presented constants are calculated by

$$\begin{cases} a_{N+1} &= \frac{r_{N-1}[0]r_N[0]}{r_{N-2}[0]r_{N-1}[1]-r_{N-1}[0]r_{N-2}[1]-r_{N-2}[0]r_N[0]}, \\ b_{N+1} &= -a_{N+1}\frac{r_{N-2}[0]}{r_{N-1}[0]}, \\ c_{N+1} &1-b_{N+1}. \end{cases}$$

Finally, there is only a missing object to calculate: the residue  $r_{N+1}$ . Its terms are obtained by

$$r_{N+1}[k] = a_{N+1}r_{N-2}[k+2] + b_{N+1}r_{N-1}[k+2] + c_{N+1}r_N[k+1].$$
(41)

The calculation stops once there are no coefficients left to use. In fact, they become shorter and shorter at each order. To summarize the whole calculation process, first of all the polynomials have to be defined thanks to (35), (36) and (37). The residues are initialized by means of (38). Then, (40) is used so that the new polynomials can be found. It is relevant to note that (35) must be employed at any order. The last step consists in obtaining the residues with (41). The process is repeated until there are no more coefficients to obtain, and as such, it is expected that the new polynomials, which obey (39), are meant to result in the best approximation. Equation (33) eventually provides the voltage solutions.

#### B. Results

Thévenin approximants are mainly used to plot PV and QV curves since they can reproduce with adequate precision both stable and unstable branches around the voltage collapse point. To show their usage, the IEEE 30 bus system has also been chosen. Active power at bus 30 has been increased at its maximum. Fig. 3 displays the profile close to the voltage collapse point. Both the positive and negative solutions of (33) have been selected.

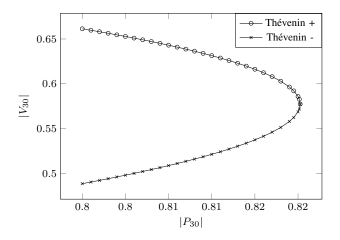


Fig. 3. PV curve around the voltage collapse point for bus number 30 of the IEEE30.

Both stable and unstable voltages are plotted with precision and they practically coincide at the voltage collapse point. Padé approximants have been found no to be an adequate tool to compute the final solution around that point because of the fact that they deviate [5]. Only the combination of them alongside with the Padé-Weierstrass process is able to match such values, although the unstable branch can not be obtained from that. This is where the advantage of using Thévenin approximants lies.

In order to validate that tool with the alternative embedding, the values of  $\left|V_{30}\right|$  obtained thanks to Thévenin approximants are compared to the ones extracted from the canonical embedding. Table III shows the absolute and relative differences. In each case the differences are extremely small. Certainly, they tend to increase when the active power is more extreme. Nevertheless, the results are satisfactory in the sense that the values obtained with both embeddings are considerably close,

TABLE III  ${\it Errors~of~|V_{30}|~obtained~with~Th\'evenin~approximants~between } \\ {\it both~embeddings}$ 

	Absolute error		Relative error	
$P_{30}$	Stable	Unstable	Stable	Unstable
-0.800	4.45E-13	3.53E-09	6.73E-13	7.22E-09
-0.805	3.93E-13	9.28E-10	6.02E-13	1.86E-09
-0.810	2.76E-13	9.38E-11	4.29E-13	1.84E-10
-0.815	7.32E-13	1.44E-10	1.16E-12	2.76E-10
-0.820	2.47E-12	1.79E-10	4.01E-12	3.33E-10
-0.825	5.74E-10	1.37E-09	9.80E-10	2.40E-09

and as a consequence, the modifications needed are assumed to be correct.

#### VI. CONCLUSION

The alternative embedding has been shown to be an appropriate approach in order to embed the equations that take part in the Holomorphic Embedding Load-Flow Method. In contrast to the canonical embedding, minor changes lead to not being required to split admittance matrices in two because of transformer tap changers as well as building an arguably more simple algorithm.

In the 11-bus ill-conditioned grid in which there are several tap changers, since the alternative embedding does not force the value of the first coefficients, it is able to reach a decent solution with substantially fewer terms than the canonical embedding.

Besides, the alternative embedding has been adapted to some ground-breaking tools such as Sigma and Thévenin approximants, which respectively help to diagnose the system and plot PV curves. The adjustments follow the same principles as in the case of the canonical embedding and they do not alter much the calculation procedure as a whole. Both tools have been successfully tested for the IEEE 30-bus system.

# ACKNOWLEDGMENT

The authors would like to thank Santiago Peñate Vera, GridCal's creator, for his help at testing the algorithm.

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