Local Network Identifiability with Partial Excitation and Measurement

Antoine Legat and Julien M. Hendrickx

Abstract—This work focuses on the identifiability of dynamical networks with partial excitation and measurement: a set of nodes are interconnected by unknown transfer functions according to a known topology, some nodes, subject to external excitation, and some nodes are measured. The goal is to determine which transfer functions in the network can be recovered based on the input-output data collected from the excited and measured nodes.

We propose a local version of network identifiability, representing the ability to recover transfer functions which are approximately known, or to recover them up to a small ambiguity. We show that local identifiability is a generic property, establish a necessary and sufficient condition in terms of matrix generic ranks, and exploit this condition to develop an algorithm determining, with probability 1, which transfer functions are locally identifiable. Our implementation presents the results graphically, and is publicly available.

I. INTRODUCTION

This paper addresses the identifiability of dynamical networks in which the node signals are connected by causal linear time-invariant transfer functions, and can be excited and/or measured. Such network can be modeled as a directed graph where each edge carries a transfer function, and known excitations and measurements are applied at certain nodes.

We consider the identifiability of a network matrix $G(z)$, where the network is made up of $n$ nodes, with node signals $w(t)$, external excitation signals $r(t)$, measured nodes $y(t)$ and noise $v_1(t), v_2(t)$ related to each other by:

$$
\begin{align*}
w(t) &= G(z)w(t) + Br(t) + v_1(t) \\
y(t) &= Cw(t) + v_2(t),
\end{align*}
$$

(1)

where matrices $B$ and $C$ are binary selections defining respectively which nodes are excited and measured, forming sets $B$ and $C$ respectively. Matrix $B$ is full column rank and each column contains one 1 and $n - 1$ zeros. Matrix $C$ is full row rank and each row contains one 1 and $n - 1$ zeros. The nonzero entries of the network matrix $G(z)$ define the network topology, and are the transfer functions to identify, forming the set of edges $E$.

We assume that the input-output relations between the excitations $r$ and measures $y$ have been identified, and that the network topology is known. From this knowledge, we aim at recovering the nonzero entries of $G(z)$.

The model (1) has recently been the object of a significant research effort. If the whole network is to be recovered, the notion of network identifiability is used, as developed in [1]. If one is interested in identifying a single module, topological conditions are derived in [2], [3]. In this paper, we do not consider the impact of noise signals $v_1, v_2$, but studying the influence of rank-reduced or correlated noise yields interesting results on identifiability [4], [5].

It turns out that the identifiability of the network, i.e. the ability to recover a module or the whole network from the input-output relation, is a generic notion: Either almost all transfer matrices corresponding to a given network structure are identifiable, in which case the structure is called generically identifiable, or none of them are. A number of works study generic identifiability when all nodes are excited or measured, i.e. when $B$ or $C = I$ [6], [7]. Considering the graph of the network, path-based conditions on the allocation of measurements in the case of full excitation are derived in [8]. Reformulating these conditions by means of disjoint trees in the graph, [9] develops a scalable algorithm to allocate excitations/measurements in case of full measurement/excitation. Rather than verifying the identifiability of a given excitation allocation, [10] addresses the question of where to allocate excitation signals under full measurement.

The conditions of [8] apply for generic identifiability, i.e. identifiability of almost all transfer matrices corresponding to a given network structure. [11] extends the path-based conditions under full excitation to determine the identifiability for all (nonzero) transfer matrices corresponding to a given structure, and [12] provides conditions for the outgoing edges of a node, and for the whole network under the same conditions.

In all these works, the common assumption is that all nodes are either excited, or measured. In [13], this assumption is relaxed and generic identifiability with partial excitation and measurement is addressed. The authors derive necessary conditions and sufficient conditions for identifying the outgoing and incoming edges of a node, and for particular network topologies such as loops and trees.

The question of generic identifiability under partial excitation and measurement for the general case remains unsolved: We wish to find a combinatorial characterization for generic identifiability of an edge or a network, that is expressed purely in terms of graph theoretical properties, akin to what was done in the full excitation case e.g. in [8]. Such characterization would in particular pave the way for optimizing the selection of nodes to be excited and measured, akin to the work in [10] in the full measurement case.
In this paper, we introduce the notion of local identifiability, i.e. we address the question of identifiability only on a neighborhood of $G(z)$. Local identifiability is a necessary condition for identifiability, and we believe its understanding will allow making progress towards understanding identifiability. We show that it is generic, provide an algebraic necessary and sufficient condition for local identifiability of a network in terms of the rank of a matrix, and a necessary and sufficient condition for local identifiability of an edge depending on the orthogonality of a matrix kernel to a certain vector (which is itself equivalent to some rank conditions). These conditions generalize previous results obtained under full excitation or measurement [8]. We then exploit these conditions to develop an algorithm that determines which transfer functions are locally identifiable, with probability 1. An implementation is available at [14], and includes a graphical representation of the results.

**Assumptions:** We consider the problem modeled in (1). Consistently with previous works, we assume the invertibility of the matrix $(I - G(z))$, which is equivalent to the network being well-posed, and that $CT(z) = C(I - G(z))^{-1}B$ has been identified exactly. We do not suppose having access to any information related to the effect of the noise signals $v_1, v_2$. The additional information that could be gathered from this knowledge in our context is left for further works.

For the simplicity of exposition, we consider in this paper a single frequency $z$, so that all transfer functions are modeled simply by a complex value, and the matrices $G$ and $T = (I - G)^{-1}$ are complex matrices rather than matrices of transfer functions. Conceptually, our generic results directly extend to the transfer function case: if one can recover a $G_{ij}(z)$ at a given frequency $z$ for almost all $G$ consistent with a network, then one can also recover it at all other frequencies, and hence recover the transfer function. We intend to remove this simplification or to formalize this intuitive argument in a further version of this work. In the remainder of this document, we omit $(z)$ to lighten notations.

**II. LOCAL IDENTIFIABILITY**

We start by reminding the definition of identifiability, see e.g. [8], [2]. A network matrix $G$ defines a directed graph where the edge $j \to i$ is present if $G_{ij} \neq 0$. We say that a matrix $G$ is consistent with a graph defined by a network matrix $G$ if $G_{ij}(z)$ is zero when there is no edge $j \to i$ in the graph defined by $G$.

**Definition 1:** The transfer function $G_{ij}$ is identifiable at $G$ from excitations $B$ and measurements $C$ if, for all network matrix $G$ consistent with the graph, there holds

$$C(I - \tilde{G})^{-1}B = C(I - G)^{-1}B \Rightarrow \tilde{G}_{ij} = G_{ij}. \quad (2)$$

The network matrix is identifiable at $G$ if each transfer function $G_{ij}$ is identifiable at $G$, i.e. if the left-hand side of (2) implies $\tilde{G} = G$.

These definitions were made for matrices of transfer functions $G(z)$, but we remind that here we consider a single frequency for simplicity, so $G$ is just a matrix in $\mathbb{C}^{n \times n}$.

The combinatorial conditions for identifiability obtained under full excitation or measurement (i.e. when $B$ or $C = I$) in [15], [6], [7], [8], [9] were made possible by a linear algebraic reformulation of the identifiability questions. No equivalent reformulation is available for settings with partial measurements and excitation. Hence we will work on a weaker notion amenable to linear analysis: local identifiability, which corresponds to identifiability provided that $\tilde{G}$ is sufficiently close to $G$.

**Definition 2:** The transfer function $G_{ij}$ is locally identifiable at $G$ if each transfer function $G_{ij}$ is locally identifiable at $G$, i.e. if the left-hand side of (3) implies $\tilde{G} = G$.

Local identifiability is a necessary condition for identifiability, both for recovering the entire network or just a particular edge. It is a priori a weaker notion, but we have so far found no example of network that is locally identifiable but not globally identifiable. Moreover, one can show that if a network is locally identifiable, then it can be recovered up to a discrete ambiguity, i.e. the set of unknown $G$ corresponding to a measured $C(I - G)^{-1}B$ would be discrete. Finally, local identifiability is relevant in the many practical situations where the transfer functions are already approximately known.

**Genericity:** Given a graph and sets $B, C$ of excited and measured nodes, we say that an edge is generically (locally) identifiable if it is (locally) identifiable at all $G$ consistent with the graph, except possibly those lying on a lower-dimensional set (i.e. a set of dimension lower than $n$). In the remainder of this paper, we will say that a property holds generically, or for almost all (resp. no) variables if it holds for all (resp. no) variables, except possibly those lying on a lower-dimensional set. We will see that local identifiability is indeed a generic notion: for given network topology and sets of excited and measured nodes, the transfer function corresponding to the edge $(i, j)$ is either identifiable at almost all $G$ (generically identifiable) or identifiable at almost none of them. More details about this will be given in future versions of this work.

**III. IDENTIFIABILITY AND INJECTIVITY**

The identifiability questions we consider can be reformulated in terms of (local) injectivity. Let $x \in \mathbb{C}^{|E|}$ be a vector compiling the $|E|$ (potentially) nonzero entries of $G$, and $G(x)$ the network matrix $G$ corresponding to a vector $x$, so
that every matrix consistent with the graph can be written as $G(x)$ for some $x$. We define

$$h : C|E| ∩ D! C|B|·|C| : x \rightarrow \text{vec} \left( (I - G(x))^{-1} B \right)$$

(4)

where the set $D \triangleq \{ x | \det(I - G(x)) = 0 \}$ collects the vectors for which the inverse does not exist. The $|B| \times |C|$ image is vectorized into a single $|B| \cdot |C|$ vector, which will ease the differential analysis below.

Recovering a matrix $G$ from $(I - G)^{-1} B$ can then be seen as recovering $x$ from $h(x)$, while recovering $G_{ij}$ corresponds to recovering a specific entry $x_{ij}$ of $x$ from $h(x)$ (where $i$ is the index of $G_{ij}$ in $x$). Local identifiability becomes then local injectivity: determining if $h(\hat{x}) = h(x)$ implies $\hat{x} = x$ for every $\hat{x}$ in a neighborhood of $x$. The local identifiability of an edge $e$ becomes a question of local coordinate-injectivity: determining if $h(\hat{x}) = h(x)$ implies $\hat{x}_e = x_e$ for every $\hat{x}$ in a neighborhood of $x$. The notion of local coordinate-injectivity is formally introduced in the definition below.

**Definition 3:** The function $f : M \rightarrow N$ is locally coordinate-injective for the coordinate $e$ at $x$ if there exists $\varepsilon > 0$ such that for all $\hat{x} \in B(x, \varepsilon)$, there holds

$$f(\hat{x}) = f(x) \Rightarrow \hat{x}_e = x_e.$$  

(5)

Intuitively, these local questions could be determined by the gradient: $h$ would be locally injective at $x$ if $\nabla h(x) \cdot \delta \neq 0$ for any nonzero $\delta \in C|E|$, i.e. if $\nabla h(x)$ has a trivial kernel. Observe this requires the number $|B| \cdot |C|$ of available input-output relations to be at least as large as the number $|E|$ of unknowns. Similarly, $h$ would be coordinate-injective at $x$ for edge $e$ if $\nabla h(x) \cdot \delta \neq 0$ for any $\delta \in C|E|$ with $\delta_e \neq 0$, i.e. if the kernel of $\nabla h(x)$ is orthogonal to $e$, the standard basis vector filled with zeros except 1 at the $e$-th entry.

However, the issue is more complex than it first seems. For example, the real function $g(x) = x^3$ is injective, but $g'(0) = 0$ admits a non-trivial kernel, in contradiction with our previous statement. The problem is that the derivative of $g$ vanishes at 0, while being nonzero elsewhere. Adding an assumption on the conservation of information contained in the derivative (i.e. the rank of its gradient) resolves that issue. The next lemma, proved in the Appendix, shows that the intuitive idea behind the gradient is in fact valid when $\nabla f$ has constant rank. We will then see in Proposition 3.1 how this rather constraining assumption can be removed provided one is only interested in local injectivity at almost all $x$.

**Lemma 3.1:** Suppose $f : M \rightarrow N$ is $C^\infty$ where $M$ and $N$ are smooth manifolds, $M$ or $N$ has finite dimension, and $\nabla f(x)$ has the same rank for every $x \in M$. Then $f$ is locally coordinate-injective for $e$ at all $x \in M$ if and only if for all $x \in M$,

$$\ker \nabla f(x) \bot e_e,$$

(6)

where $e_e$ denotes the standard basis vector filled with zeros except 1 at the $e$-th entry.

The next proposition, proved in the Appendix, characterizes local injectivity almost everywhere for analytic functions. Analytic functions are functions which can be locally approximated everywhere by their Taylor expansion, see e.g. [16] for a precise definition. They include in particular all compositions of products, sums, divisions, etc.

**Proposition 3.1:** Let $f : M \rightarrow N$ be an analytic function where $M$ and $N$ are smooth manifolds, $M$ or $N$ has finite dimension, and $M$ is open. Then exactly one of the two following holds:

(i) $\ker \nabla f(x) \bot e_e$ for almost all $x$ and $f$ is locally coordinate-injective for $e$ at almost all $x$;

(ii) $\ker \nabla f(x) \bot e_e$ for almost no $x$ and $f$ is locally coordinate-injective for $e$ at almost no $x$.

**IV. NECESSARY AND SUFFICIENT CONDITIONS FOR LOCAL IDENTIFIABILITY**

First we define the $|B| \cdot |C| \times |E|$ matrix $K$ which, as will be proved in Lemma 4.1, is the gradient of $h$.

$$K(x) \triangleq (B^T T^T(x) \otimes CT(x)) I_G,$$

(7)

where $T(x) \triangleq (I - G(x))^{-1}$, $\otimes$ denotes the Kronecker product and the matrix $I_G \in \{0,1\}^{n^2 \times |E|}$ selects only the columns of the preceding $|B| \cdot |C| \times n^2$ matrix corresponding to actual edges. It is defined by

$$I_G \triangleq \left[ \text{vec}(G(e_1)) \ \text{vec}(G(e_2)) \ \cdots \ \text{vec}(G(e_{|E|})) \right].$$

We are now ready to show that local identifiability is generic, and derive our necessary and sufficient condition.

**Theorem 4.1:** Let $K$ be as defined in (7) and $e$ be the index of $G_{ij}$ in $x$. Exactly one of the two following holds:

(i) $\ker K(x) \bot e_e$ for almost all $x$ and $G_{ij}$ is generically locally identifiable;

(ii) $\ker K(x) \bot e_e$ for almost no $x$ and $G_{ij}$ is generically locally non-identifiable, hence generically non-identifiable.

Moreover, $\ker K(x) \bot e_e$ is equivalent to the following implication holding for all $\Delta$ consistent with the graph:

$$(I - G)^{-1} \Delta (I - G)^{-1} B = 0 \Rightarrow \Delta_{ij} = 0.$$  

(8)

The proof of Theorem 4.1 relies on applying Proposition 3.1 on the gradient of $h$, characterized by Lemma 4.1.

**Lemma 4.1:** Let $h$ be the function defined in (4). Its gradient is given by

$$\nabla h(x) = (B^T T^T(x) \otimes CT(x)) I_G = K(x).$$

(9)

Besides, for all $\delta \in C|E|$, there holds

$$\text{mat} (\nabla h(x) \delta) = CT(x) G(\delta) T(x) B,$$

(10)

where $\text{mat}$ reorganizes its $|B| \cdot |C|$ vector in a $|B| \times |C|$ matrix.
Proof: First note that \( \nabla h(x) \delta \) is the differential of \( h \) at \( x \) evaluated on \( \delta \), which we denote \( d_x h(\delta) \). We start by deriving the expression of the differential (10), from which we compute the gradient (9). The differential of \( h \) follows from the differential of \( T \) by linearity: \( \text{mat}(dh) = C dTB \), which relies on its partial derivatives, developed as [17]:

\[
\frac{dT}{dx_e}(x) = - (I - G(x))^{-1} \frac{\partial (I - G)}{\partial x_e} (I - G(x))^{-1}
\]

\[
= -T(x) \left( \frac{\partial G}{\partial x_e}(x) \right) T(x) = T(x) G(e_e) T(x).
\]

The partial derivatives then yield the differential:

\[
d_x T(\delta) = \sum_{e=1}^{\left|E\right|} T(x) G(e_e) T(x) \delta_e = T(x) G(\delta) T(x).
\]

For the differential of \( h \), we obtain

\[
\text{mat}(d_x h(\delta)) = C d_x T(\delta) B = CT(x) G(\delta) T(x) B.
\]

In order to derive the gradient of \( h \), we vectorize (11):

\[
\nabla h(x) \delta = \text{vec} \left( CT(x) G(\delta) T(x) B \right)
\]

\[
= \left( B^T T^T(x) \otimes CT(x) \right) \text{vec} \left( G(\delta) \right),
\]

where vec stacks the columns of its \( |B| \times |C| \) matrix argument into a \( |B| \times |C| \) vector. Then, observing that \( \text{vec} \left( G(\delta) \right) = I_G \delta \) gives \( \nabla h(x) \delta = K(x) \delta \ \forall \delta \in \mathbb{C}^{|E|} \), which yields (9).

**Proof of Theorem 4.1:** As explained in Section III, the generic local identifiability of transfer function \( G_{ij} \) is equivalent to function \( h \) defined in (4) being locally coordinate-injective for \( e \) at almost all \( x \) (\( e \) is the index of \( G_{ij} \) in \( x \)).

Proposition 3.1 gives conditions on local coordinate-injectivity, let us verify its assumptions for \( h \). First \( h \) is analytic, and its domain and image sets have finite dimension. The openness of the domain \( \mathbb{C}^{|B|} \setminus \mathcal{D} \) follows from the continuity of \( \det(I - G) \). Since the inverse image of a closed set by a continuous function is a closed set [18], the zero set of \( \det(I - G) \) is closed, and its complement is open. Its assumptions being fulfilled, Proposition 3.1 applies. The first part of the theorem then follows directly from (9).

Besides, \( \ker \nabla h(x) \perp e_e \) can be rewritten equivalently as

\[
\nabla h(x) \delta = 0 \Rightarrow \delta_e = 0.
\]

Reorganizing this \( |B| \times |C| \) vector into a \( |B| \times |C| \) matrix gives

\[
\text{mat} \left( \nabla h(x) \delta \right) = 0 \Rightarrow \delta_e = 0,
\]

and combining it with (10) then yields (8).

Theorem 4.1 provides a necessary and sufficient condition for local identifiability of a transfer function. Therefore, combining this condition for all edges immediately leads to a characterization of local identifiability of the whole network.

**Corollary 4.1:** Exactly one of the two following holds:

(i) \( \text{rank} K(x) = |E| \) for almost all \( x \) and \( G \) is generically locally identifiable;

(ii) \( \text{rank} K(x) = |E| \) for almost no \( x \) and \( G \) is generically locally non-identifiable.

Moreover, \( \text{rank} K(x) = |E| \) is equivalent to the following implication holding for all \( \Delta \) consistent with the graph:

\[
C(I - G)^{-1} \Delta (I - G)^{-1} B = 0 \Rightarrow \Delta = 0. \quad (12)
\]

Proof: First, let \( K(x) \perp e_e \) for each \( e \) is equivalent to \( \ker K(x) = \{0\} \), and therefore \( \text{rank} K(x) = |E| \) by the rank-nullity theorem. Then, (8) for each \((i,j)\) yields (12).

We observe that condition (12) reduces to the necessary and sufficient condition for (global) identifiability in the full excitation (resp. measurement) case. Indeed, since \( T = (I - G)^{-1} \) is by construction invertible, \( CT\Delta B = 0 \) is equivalent to \( CT\Delta = 0 \) when \( B \) is the identity matrix (resp. to \( \Delta B \) when \( C \) is the identity matrix), as in [8].

**V. ALGORITHM**

The necessary and sufficient conditions of Theorem 4.1 can be tested exactly by symbolic computation, but it becomes rapidly computationally intractable. Hence, we have designed a probability-1 algorithm that exploits the generic character of local identifiability, as proved in Theorem 4.1. Testing these conditions with a randomly selected \( x \in \mathbb{C}^{|E|} \) will therefore provide the correct result with a probability 1, and can be done very efficiently. This extends what E. J. Davison does in the controllability context [19].

Although this idea allows to design an algorithm working with probability 1, we cannot exclude the possibility that an actual implementation could suffer from numerical issues for large networks, with e.g. the numerically computed rank being very close to zero if the randomly selected values are close to the problematic lower-dimensional set. Hence we repeat the procedure several times to increase the reliability of our results, as described in Algorithm 1. However, we did not observe any such numerical problem in any of our tests. Our implementation is available at [14], and includes a graphical representation of the results.

Figure 1 shows examples of outputs of Algorithm 1. Figure 1(a) is the network of Example 2 in [9], for which the authors provide an optimal excitation set assuming all nodes are measured. Our algorithm shows that all edges can still be locally identified if some of the nodes (i.e., 4, 10, 11) are not actually measured, and node 9 must not be excited. Note this does not contradict the optimality of the excited set found in [9], since this set was found under the assumption of full measurement, and node 9 was affected by external noise.

Figure 1(b) shows a nontrivial example of network where all edges but two are generically locally identifiable. Interestingly, exciting 1 or 9 allows recovering both (1, 11) and (9, 6), even though these edges do not appear directly related.
Algorithm 1 Identifiability test

Require: Graph topology, matrices $B$ and $C$, $n_{\text{samples}}$
Result: The identifiability of each transfer function
1: Initialize network to false and edges to a false
2: for $i \leftarrow 1$ to $n_{\text{samples}}$ by 1 do
3: Randomly generate a complex network matrix $G$
4: Construct $K = (B^T T^T \otimes CT) I_G$
5: if rank $K = |E|$ then
6: network $\leftarrow$ true
7: else Compute $V$, basis of ker $K$
8: Calculate $v$, the binary vector of length $|E|$, with $v_e = \text{true}$ if the $e$-th entry of each vector of $V$ is 0, $v_e = \text{false}$ otherwise
9: edges $\leftarrow$ edges or $v$ entry-wise
10: if network then return Network identifiable
11: else return edges

VI. CONCLUSION

This work was motivated by one main open question: determining graph-theoretical or combinatorial conditions for generic network and edge identifiability in networked systems.

The local notions we have introduced allowed making progress in this direction. In particular, the necessary and sufficient conditions for generic local identifiability in terms of generic ranks allowed us to design an algorithm determining, with probability 1, the generic local identifiability of each edge.

Providing a graph-theoretical characterization for local identifiability remains an open question, but we believe that our generic rank-based characterization could pave the way to a solution. Moreover, our algorithm allows rapidly testing ideas and conjectures, hence facilitates further research.

In principle, generic local identifiability is a weaker notion than generic identifiability. However, we were so far unable to find examples of networks that are locally, but not globally, identifiable. The possible equivalence of the two notions would greatly simplify the study of identifiability, and remains an open question.

VII. ACKNOWLEDGMENTS

The authors gratefully acknowledge P.-A. Absil for his help on manifold theory, and M. Gevers and A. S. Bazanella for the interesting and insightful discussions.

REFERENCES

[1] H. H. Weerts, A. G. Dankers, and P. M. Van den Hof, “Identifiability in dynamic network identification,” IFAC-PapersOnLine, vol. 48, no. 28, pp. 1409–1414, 2015.
[2] H. Weerts, P. M. Van den Hof, and A. Dankers, “Single module identifiability in linear dynamic networks,” in 2018 IEEE Conference on Decision and Control (CDC), pp. 4725–4730, IEEE, 2018.
[3] M. Gevers, A. S. Bazanella, and G. V. da Silva, “A practical method for the consistent identification of a module in a dynamical network,” IFAC-PapersOnLine, vol. 51, no. 15, pp. 862–867, 2018.
[4] H. H. Weerts, P. M. Van den Hof, and A. G. Dankers, “Prediction error identification of linear dynamic networks with rank-reduced noise,” Automatica, vol. 98, pp. 256–268, 2018.
[5] P. M. Van den Hof, K. R. Ramaswamy, A. G. Dankers, and G. Bottegal, “Local module identification in dynamic networks with correlated noise: the full input case,” in 2019 IEEE 58th Conference on Decision and Control (CDC), pp. 5494–5499, IEEE, 2019.
[6] A. S. Bazanella, M. Gevers, J. M. Hendrickx, and A. Parraga, “Identifiability of dynamical networks: which nodes need to be measured?” in 2017 IEEE 56th Annual Conference on Decision and Control (CDC), pp. 5870–5875, IEEE, 2017.
[7] H. H. Weerts, P. M. Van den Hof, and A. G. Dankers, “Identifiability of linear dynamic networks,” Automatica, vol. 89, pp. 247–258, 2018.
[8] J. M. Hendrickx, M. Gevers, and A. S. Bazanella, “Identifiability of dynamical networks with partial node measurements,” IEEE Transactions on Automatic Control, vol. 64, no. 6, pp. 2240–2253, 2018.
[9] X. Cheng, S. Shi, and P. M. Van den Hof, “Allocation of excitation signals for generic identifiability of dynamic networks,” in Proceedings of the IEEE Conference on Decision and Control, 2019.
[10] S. Shi, X. Cheng, and P. M. Van den Hof, “Excitation allocation for generic identifiability of a single module in dynamic networks: A graphic approach,” 2019.
[11] H. J. van Waarde, P. Tesi, and M. K. Camlibel, “Topological conditions for identifiability of dynamical networks with partial node measurements,” IFAC-PapersOnLine, vol. 51, no. 23, pp. 319–324, 2018.
[12] H. J. Van Waarde, P. Tesi, and M. K. Camlibel, “Necessary and sufficient topological conditions for identifiability of dynamical networks,” IEEE Transactions on Automatic Control, 2019.
A. S. Bazanella, M. Gevers, and J. M. Hendrickx, “Network identification with partial excitation and measurement,” in 2019 IEEE 58th Conference on Decision and Control (CDC), pp. 5500–5506, IEEE, 2019.

A. Legat and J. M. Hendrickx, “Identifiability test. https://github.com/alegat/identifiable.”

M. Gevers, A. S. Bazanella, and A. Parraga, “On the identifiability of dynamical networks,” IFAC-PapersOnLine, vol. 50, no. 1, pp. 10580–10585, 2017.

E. Hille, Analytic function theory, vol. 2. American Mathematical Soc., 2005.

K. Petersen and M. Pedersen, “The matrix cookbook, version 20121115,” Technical Univ. Denmark, Kongens Lyngby, Denmark, Tech. Rep., vol. 3274, 2012.

F. Hausdorff, Set theory. American Mathematical Society (RI), 1991.

E. J. Davison, “Connectability and structural controllability of composite systems,” Automatica, vol. 13, no. 2, pp. 109–123, 1977.

R. Abraham, J. E. Marsden, and T. Ratiu, Manifolds, tensor analysis, and applications, vol. 75. Springer Science & Business Media, 2012.

J. Van der Woude, “A graph-theoretic characterization for the rank of the transfer matrix of a structured system,” Mathematics of Control, Signals and Systems, vol. 4, no. 1, pp. 33–40, 1991.

J. P. D’Angelo, An introduction to complex analysis and geometry, vol. 12. American Mathematical Soc., 2010.

APPENDIX

A. Proof of Lemma 3.1

We must prove that (5) for all \( x \in M \Leftrightarrow (6) \) for all \( x \in M \). The smoothness of \( f \) and constant rank of \( \nabla f \) allow the application of the submersion theorem [20]. In short, this theorem states that the level set of an image \( n_0 \) by such function is a submanifold of \( M \), and its tangent space coincides with the kernel of the gradient. Here, we take \( n_0 = f(\hat{x}) \) to make the connection with injectivity, and the theorem yields that for all \( x \), \( f^{-1}(f(x)) \) is a submanifold of \( M \) with

\[
T_{\hat{x}}f^{-1}(f(x)) = \ker \nabla f(\hat{x}) \ \forall \ \hat{x} \in f^{-1}(f(x)),
\]

where \( T_{\hat{x}}A \) denotes the tangent space to manifold \( A \) at \( \hat{x} \).

We start with necessity: assume (5) for all \( x \), and consider a specific \( x \). Each \( \hat{x} \) belonging to a neighborhood of \( x \) in \( f^{-1}(f(x)) \) has the same \( e \)-coordinate as \( x \). Therefore, any tangent vector to \( f^{-1}(f(x)) \) at \( x \) has a zero \( e \)-component, and the tangent space to this manifold at \( x \) is orthogonal to direction \( e \), i.e. \( T_{\hat{x}}f^{-1}(f(x)) \perp e \). Equation (13) evaluated at \( \hat{x} = x \) gives (6) and proves necessity.

We now prove sufficiency: assume (6) for all \( x \), and consider a specific \( x \). Combining (6) with (13) yields \( T_{\hat{x}}f^{-1}(f(x)) \perp e \), for all \( \hat{x} \in f^{-1}(f(x)) \). Take a specific \( \hat{x} \in f^{-1}(f(x)) \), and assume that there is a path lying in \( f^{-1}(f(x)) \) between \( x \) and \( \hat{x} \). At every point of this path, the tangent space is orthogonal to direction \( e \), so the derivative along the path is zero in the \( e \)-coordinate. Integrating along the path gives a zero increase in \( e \)-coordinate, hence \( \hat{x}_e = x_e \).

To complete the proof, we need to show that there exists \( \epsilon > 0 \) such that for all \( \hat{x} \in B(x, \epsilon) \cap f^{-1}(f(x)) \), there is a path lying in \( f^{-1}(f(x)) \) between \( x \) and \( \hat{x} \). Such path can be found with the rank theorem [20]. Roughly speaking, this theorem states that if \( \nabla f \) has constant rank \( k \) in a manifold of dimension \( m \), then \( m - k \) variables are redundant and can be eliminated. More precisely, for all \( p \) in the manifold, there exist local coordinates centered at \( p \) and \( f(p) \) in which \( f(q) = (q_1, \ldots, q_k, 0, \ldots, 0) \) for any \( q \) in the manifold.

For the rest of this proof, we work in the local coordinates provided by this theorem, centered at \( p = x \). Since these are centered at \( x \) and \( f(x) \), \( x_1 = \cdots = x_k = 0 \). Let us now take \( \epsilon > 0 \), and \( \hat{x} \in B(x, \epsilon) \cap f^{-1}(f(x)) \). The rank theorem yields that the last entries of \( f(\hat{x}) \) are zero in those local coordinates, i.e. \( f(\hat{x}) = (\hat{x}_1, \ldots, \hat{x}_k, 0, \ldots, 0) \). Since \( \hat{x} \) lies in the manifold \( f^{-1}(f(x)) \), it must have the same image as \( x \), hence \( f(\hat{x}) = f(x) = 0 \), and \( \hat{x}_1 = \cdots = \hat{x}_k = 0 \). An example of path between \( x \) and \( \hat{x} \) is given by \( (0, \ldots, 0, \lambda \hat{x}_{k+1}, \ldots, \lambda \hat{x}_{|E|}) \), with \( \lambda \in [0, 1] \). Applying the function \( f \) on any point of this path gives 0 in the local coordinates by the rank theorem, showing that the path is indeed included in the manifold \( f^{-1}(f(x)) \).

B. Proof of Proposition 3.1

The proof of Proposition 3.1 relies on Lemma B.1.

**Lemma B.1:** The gradient of an analytic function \( f \) reaches its maximal rank everywhere except on a closed lower-dimensional set. Moreover, the orthogonality relation

\[
\ker \nabla f(x) \perp e_e
\]

either holds:

(i) for all \( x \) except those lying on a closed lower-d. set;
(ii) only for \( x \) lying on a closed lower-dimensional set.

**Sketch of proof:** First, the set on which the rank of a matrix drops can be expressed as the intersection of zero sets of determinants of submatrices [21]. The determinant is analytic in its entries, which are analytic in \( x \) since \( f \) is assumed analytic (hence so is \( \nabla f \)). Since nonconstant analytic functions vanish only on a lower-dimensional set [22], the set on which \( \nabla f(x) \) does not reach its maximal rank has lower dimension. Besides, this set is also closed: the inverse image of a closed set by a continuous function is a closed set [18], hence the zero sets of the determinants are closed, and so is their intersection. This extends the notion of generic rank introduced in [19].

Then, (14) can be equivalently rewritten as \( \nabla f(x)\delta = 0 \Rightarrow \delta_e = 0 \ \forall \ \delta \), which is equivalent to the linear independence of the \( e \)-th column of \( \nabla f(x) \) with the other columns. It can be formulated by means of ranks, and a reasoning similar to above completes the proof.

**Proof of Proposition 3.1:** We denote \( \tilde{M} \) the intersection of the set on which \( \nabla f \) reaches its maximal rank (hence constant), and the set on which (14) holds (case (i)) or does not hold (case (ii)). Lemma B.1 ensures that those two sets cover the whole domain \( M \) except a closed lower-dimensional set. Hence, so does \( \tilde{M} \), and it is open since \( M \) is assumed open. Thus, there is an \( \epsilon \) for every \( x \in M \) such that the rank of \( \nabla f \) is constant over \( B(x, \epsilon) \), and (14) holds (case (i)) or does not hold (case (ii)) on the ball. Denote \( f_{x,\epsilon} \) the restriction of \( f \) on \( B(x, \epsilon) \). The result follows by applying Lemma 3.1 on \( f_{x,\epsilon} \) for all \( x \in \tilde{M} \).