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Modular response analysis reformulated as a multilinear regression problem

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Abstract

\textbf{Motivation:} Modular response analysis (MRA) is a well-established method to infer biological networks from perturbation data. Classically, MRA requires the solution of a linear system, and results are sensitive to noise in the data and perturbation intensities. Due to noise propagation, applications to networks of 10 nodes or more are difficult.

\textbf{Results:} We propose a new formulation of MRA as a multilinear regression problem. This enables to integrate all the replicates and potential additional perturbations in a larger, over-determined, and more stable system of equations. More relevant confidence intervals on network parameters can be obtained, and we show competitive performance for networks of size up to 1000. Prior knowledge integration in the form of known null edges further improves these results.

\textbf{Availability and implementation:} The R code used to obtain the presented results is available from GitHub: https://github.com/J-P-Borg/BioInformatics

1 Introduction

Biological systems are orchestrated by a multitude of interaction networks. At a cellular scale, the knowledge of protein physical interaction networks, phosphorylation cascades, or gene regulatory networks is central in our understanding of homeostasis, disease, reaction to environmental changes, and many other situations. The inference of biological networks from experimental data has thus attracted much attention from the computational biology community (Huynh-Thu and Sanguinetti 2019).

A common experimental design to learn about the connectivity between a set of molecules of interest is to measure their activity under different conditions. Molecules can be for instance genes, transcripts, proteins, or metabolites, and the notion of activity might relate to their abundance or state, e.g. the phosphorylation level of a protein. Modular response analysis (MRA) is a versatile framework allowing to infer connectivity between molecules (or groups of molecules called modules) $X_1, \ldots, X_N$ simply assuming that they are related by an equation of the form $dX/dt = f(X)$, $X = (X_1, \ldots, X_N)^T$, and a particular experimental design. This design requires that measures of $X_i$ activities are obtained under the systematic perturbation of each molecule activity, and the system finds itself in a steady state for each of those perturbations. The elegance of MRA is that the steady-state assumption enables applying the implicit function theorem leading to a first-order approximation of $f(X)$ without knowing this function explicitly (Kholodenko et al. 2002). Except for the steady-state hypothesis, which is not fulfilled by every system obviously, MRA provides a generic solution involving linear algebra only to explore biological networks quantitatively. It has been applied and extended by a number of researchers (Dorel et al. 2018; Santra et al. 2018; Jimenez-Dominguez et al. 2021; Mekedem et al. 2022). However, MRA is sensitive to measurement noise and to the intensity of the perturbations exerted on network nodes (Andrec et al. 2005; Thomaseth et al. 2018).

Different approaches have been suggested to alleviate these problems (Santra et al. 2018), e.g. to perform a bootstrap followed by an estimation of confidence intervals (CIs) of the connectivity coefficients (Santos et al. 2007; Jimenez-Dominguez et al. 2021).

In this report, we combine MRA with multilinear regression. Indeed, MRA equations can be interpreted as a linear regression problem (see below) thus enabling the use of any linear regression strategy. This new formulation hence defines a family of methods, which we will illustrate employing classical algorithms such as the least square, LASSO, or STEP. One advantage of the regression approach is to integrate the treatment of the experimental noise with the model accuracy estimation in terms of residual variance. That is,
how well the linear MRA model approximates the potentially non-linear biological system and how accurate are the estimated network parameters. Using data from DREAM 4 Challenge, we show that we can apply MRA modeling to 10- to 100-node networks with competitive performance, while a standard resolution of MRA equations already encounters difficulties with 10-node networks. Using realistic synthetic networks of sizes up to 1000 nodes (Carré et al. 2017) and varying noise in the data, we confirm those results. We further compare MRA and our multilinear regression variants against general-purpose network inference algorithms such as ARACNE (Margolin et al. 2006), CLR (Faith et al. 2007), and MRNET (Meyer et al. 2007), which we largely outperform.

2 Materials and methods

2.1 Fundamental definitions

We consider a putative biological network comprised of $N$ modules that potentially interact. A module can be a molecule (transcript, protein, . . .) or it can represent the contribution to the network of a set of molecules such as a pathway or part of a pathway. In the latter case, one measurement reports for the activity of the whole module. MRA aims at determining signed connectivity coefficients between the modules. The module activity levels are represented as a time-dependent function $X: \mathbb{R}^+ \rightarrow \mathbb{R}^N$. In case the modules were gene transcripts, $X(t)$ could for instance represent their abundance over time. We assume the existence of a continuously differentiable function $f: \mathbb{R}^N \times \mathbb{R}^M \rightarrow \mathbb{R}^N$ and a vector of intrinsic parameters $P \in \mathbb{R}^M$, at least one per module (hence $M \geq N$), such that

$$\frac{dx}{dt} = f(X, P).$$  \hfill (1)

The function $f$ is usually unknown. $P_0$ represents the biological system in its basal state, i.e. in the absence of any perturbation. We suppose the existence of a time $t_0$ after which the system has reached a steady state (including all the perturbed states):

$$\frac{dx}{dt} = 0, \forall t \geq t_0. \hfill (2)$$

Let $X = X(P_0)$ be a solution of the system (2). The application of the implicit function theorem gives us an exact formula for the connectivity coefficient representing the influence of a module $j$ on another module $i$ (Kholodenko et al. 2002; Jimenez-Dominguez et al. 2021):

$$r_{ij} = -\left( \frac{X_i}{X_j} \right) \left( \frac{\partial X_i}{\partial X_j} \right) = \frac{X_i}{X_j} \left( \frac{\partial X_i}{\partial X_j} \right), \quad 1 \leq i \neq j \leq N. \hfill (3)$$

Note that $r_{ij}$ is not defined by Equation (3). By convention, we set $r_{ii} = -1$, see Jimenez-Dominguez et al. (2021) for an algebraic justification.

Taylor’s development at the first order enables us to write an expression for the system under perturbation $\Delta P$.

$$X_i(P_0 + \Delta P) - X_i(P_0) = \sum_{j=1, j \neq i}^{N} \frac{X_i}{X_j} \left( \frac{\partial X_i}{\partial X_j} \right) (X_j(P_0 + \Delta P) - X_j(P_0))$$

$$+ \frac{1}{X_i} \sum_{j=1}^{N} \left( \frac{\partial X_i}{\partial P_j} \right) \Delta P_j + o(||\Delta P||),$$

for $1 \leq i \leq N$ and with $o(b) = 0(b)$, $\lim_{b \rightarrow 0} O(\theta) = \theta$ continuous in the vicinity of 0.

2.1.1 Standard MRA formulation

Only one perturbation per module is considered implying $M = N$. Experimental measurements obtained under perturbation $q_i$ result from the modification of the intrinsic parameter $P_i$ that directly affects node $i$ only. Accordingly,

$$\frac{\partial X_i}{\partial P_i} \neq 0 \quad \text{if} \quad i = j \quad \text{and} \quad \frac{\partial X_i}{\partial P_i} = 0 \quad \text{if} \quad i \neq j. \hfill (4)$$

We introduce the notation

$$\frac{\Delta X_i}{X_i} \bigg|_{q_i} = 2 \frac{X_i(P_0 + \Delta P_i) - X_i(P_0)}{X_i(P_0 + \Delta P_i) + X_i(P_0)} \hfill (5)$$

Using the mid-point in the denominator of Equation (6) right-term instead of $X_i(P_0)$ is customary to avoid divisions by 0. Thanks to Equations (4) and (5), we obtain the system of equations with $1 \leq i \leq N$ and $k \neq i$:

$$\frac{\Delta X_i}{X_i} \bigg|_{q_i} = \sum_{j=1, j \neq i}^{N} r_{ij} \left( \frac{\Delta X_j}{X_j} \right) \bigg|_{q_i} + \sigma(||\Delta P||). \hfill (6)$$

Ignoring the error term $\sigma(||\Delta P||)$ in (7) that represents departure from linearity, we obtain its linear approximation

$$\frac{\Delta X_i}{X_i} \bigg|_{q_i} = \sum_{j=1, j \neq i}^{N} r_{ij} \left( \frac{\Delta X_j}{X_j} \right) \bigg|_{q_i} \hfill (7)$$

that is solved by MRA (note that $r_{ij}$ replaces the exact $r_{ij}$). In particular, writing matrices $r = (r_{ij})$ and $R = \left( \frac{\Delta X_j}{X_j} \right) \bigg|_{q_i}$, the connectivity coefficients are obtained by:

$$r = -\left[ \text{diag}(R^{-1}) \right]^{-1} R^{-1} \hfill (8)$$

an $N \times N$ system (Kholodenko et al. 2002).

2.2 A multilinear regression formulation

Equation (9) does not take into account the existence of replicates in the data. It requires a single value for each perturbed state and each module. In practice, ad hoc procedures must be added to exploit replicates properly and estimate CIs for the $r_{ij}$ computed after Equation (9). Typically, bootstrap approaches are used that apply Equation (9) to resampled data or Equation (9) is applied to averaged data perturbed by an appropriate noise function (Andrec et al. 2005; Santos et al. 2007; Thomas et al. 2018; Jimenez-Dominguez et al. 2021).

An alternative approach is to consider Equation (8) as the homogeneous multilinear regression of $\frac{\Delta X_i}{X_i}$ with regression parameters $r_{ij}$. In that case, the system to solve is not limited to $N - 1$ equations. We can have one equation per replicate with any number of replicates. It is even possible to integrate multiple perturbations at each node, e.g. at different intensities or by different means, and hence $M \geq N$. The number of perturbations is also no longer required to be identical at each node. Obviously, we have $N$ such regressions to perform ($N - 1$ variables each) and the regression approach let us estimate $r_{ij}$ variability, directly taking advantage of all the tools provided by the regression literature ( Hastie et al. 2009).

2.3 Considered regression methods

To solve Equation (8) with a regression approach leaves the choice of the specific regression method free. Here, we considered several standard methods. An obvious choice was least square estimation (LSE) applied to multilinear regression. Because most biological networks are rather sparse and they can reach a certain size, we primarily considered methods able to eliminate the least significant regression parameters. Accordingly, we defined the LSE_CI method as the application of LSE followed by an elimination of all the $r_{ij}$ whose 95% CI included 0. A second method was threshold linear regression (TLR), which also starts with LSE but eliminates coefficients differently. The rule is to compute a threshold $T_h$ as the 25th percentile of all the
estimated values \(|r_j|\) and to set to 0 those \(r_j\) that are below threshold in absolute value.

The third method was LASSO, a shrinkage estimator. Using LASSO for a node \(i\) (the \(i\)th of the \(N\) systems of size \(N - 1\) to solve) is defined by:

\[
r_j^{\text{Lasso}} = \arg\min_{r_j} \left( \sum_{k=1,k \neq i}^N \left( \frac{\Delta X_{ik}}{X_i} r_k - \frac{N}{j \neq i} r_j \frac{\Delta X_{ij}}{X_i} r_k \right)^2 \right). \tag{10}
\]

The choice of the hyper-parameter \(\lambda_i\) is determined by cross-validation (CV) (Hastie et al. 2009), and it is hence necessary to have experimental replicates or multiple perturbations at certain nodes.

The fourth option was methods integrating a variable selection scheme. Namely, we used STEP forward (STEP-Fo), STEP backward (STEP-Ba), and Stepwise regression (STEP-Bo) combining both backward and forward (Hastie et al. 2009). They essentially consist in finding subsets of variables minimizing the sum of the residual squares while ignoring the other variables according to different selection strategies. This selection process introduces a bias in the solution, but here, this bias should remain modest because the networks are sparse (roughly 80% of null edges).

2.4.1 Selection and discretization of connectivity coefficients

Standard MRA and the regression-based variants we propose all estimate real-valued connectivity coefficients \(r_{ij}\). Methods such as LASSO, LSE, CI, TLR, and STEP-xx include a mechanism to identify coefficients deemed to be significant, the other ones being set to 0. MRA usually outputs coefficients \(r_{ij}\) that are all or almost all nonzero. We applied two techniques to assess standard MRA connectivity coefficient significance. For very small networks (the 6-kinase example), we used a bootstrap (100 repetitions) to estimate a CI on each \(r_{ij}\) and set \(r_{ij}\) to 0 as soon as the CI included 0. For larger networks, this procedure was too slow and we applied a heuristics: we defined a threshold \(T \approx \frac{\text{card}(0)}{\text{card}(1)}\) introduced by Kholodenko et al. (2002) in MRA original reference network after discretization, we defined \(T\approx \frac{\text{card}(0)}{\text{card}(1)}\).

In cases where it was necessary to compare with a reference network where only the existence and sign of edges were known, we applied a bootstrap (100 repetitions) to estimate a CI on each \(r_{ij}\) and set \(r_{ij}\) to 0 as soon as the CI included 0. For larger networks, this procedure was too slow and we applied a heuristics: we defined a threshold \(T \approx \frac{\text{card}(0)}{\text{card}(1)}\).

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2.4 Network inference methods

We considered ARACNE (Margolin et al. 2006), CLR (Faith et al. 2007), and MRNET (Meyer et al. 2007). We used their implementation in the Bioconductor R package minet (Meyer et al. 2008).

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2.5 Generation of synthetic networks with FRANK

FRANK (Carre et al. 2017) was used requiring one eigenvalue of the TF matrix to reside on the unit circle to ensure reaching a steady state, dynamic mode, minimum sparsity 15% (meaning that 85% of the edges are null), and the other parameters were left at their default values. The output of FRANK provided us with matrices that we used as the exact, reference matrix of connectivity coefficients \(r^e\) after setting the diagonal elements at -1 to comply with MRA formalism. To generate the input data for the inference methods, we had to compute the corresponding matrix \(R\). From Equation (9), we obtain easily that \(R = r^{-1} (\text{diag}(r^{-1}))^{-1} \text{diag}(R)\), and \(\text{diag}(R)\) is known since we applied known perturbations to \(X_i\) concentrations. For simulating full KOs we apply -100% to each \(X_i\) and after Equation (6) all the diagonal elements of \(R\) are equal to -2. For a KO at -50% they are all equal to -2/3. Using the matrix \(r^e\) from FRANK in these equations gives the corresponding \(R\). Additive \(N(0, \sigma)\) noise was added to \(R\) with \(\sigma = kX\), \(k\) a factor adjusting the noise level (see Section 3) and \(X\) the average concentration of the genes.

2.6 Prior knowledge integration

Let us assume that we know some null edges of node \(X_i\), i.e., a set \(A_i = \{j \in \{1, \ldots, N\}; r_{ij} = 0\}\).

Each index \(j \in A_i\) cancels a column of the linear system associated with solving node \(X_i\) connectivity coefficients. When the regressive approach is used, Equation (8) becomes:

\[
\frac{\Delta X_i}{X_i} = \sum_{j=1,j \neq i}^N r_{ij} \frac{\Delta X_j}{X_j} \cdot \tag{11}
\]

Hence, the linear system corresponding to node \(X_i\) is reduced to \(N - 1 - \text{card}(A_i)\) parameters to estimate. The number of degrees of freedom of the model and its residual variance decrease accordingly.

3 Results and discussion

3.1 Impact of noise and perturbation intensities on MRA solutions

Conceptually, the derivation of MRA equations entails infinitesimal perturbations of the network modules (Kholodenko et al. 2002) such that differential calculus can be applied, notably the implicit function theorem that is a local result (Jimenez-Dominguez et al. 2021). In practice, infinitesimal perturbations are usually not feasible and not advisable because experimental data tend to be noisy. Nevertheless, Taylor’s development in Equation (4) shows a dependence of the error on the perturbation intensity (error term) underlying a potential loss of accuracy of MRA depending on the strengths of the variable nonlinear dependencies. Accordingly, we decided to study the relationship between accuracy, noise, and perturbation strengths on a small but representative system.

We used the synthetic kinetics network (MAP, MKK, and MKKK) introduced by Kholodenko et al. (2002) in MRA original paper. This system is comprised of six molecules (counting the phosphorylation states) and, unlike Kholodenko’s analysis that was limited to three modules, we considered the whole system with six modules, one per molecule (Fig. 1A). The dynamical equations are given in Supplementary Information. We solved them numerically using the R package ode and determined the time \(t_0\) after which a
steady state was reached (details in Supplementary Information). The exact connectivity coefficients $r_{ij}$ (Fig. 1B) were computed numerically.

We started with data devoid of noise and perturbations $q_{kl}$ were applied to the six-speed constants of the system $\mu_{m+\Delta P}^i, i \in \{3, 4, 7, 8, 11, 12\}$ (see Supplementary Information) with $\Delta P$ equals to ±50%. Such perturbations could be induced by kinase inhibitors for instance. For each perturbation, a new steady state $X(P_i + \Delta P)$ was computed; the perturbation was applied to each node successively. For a ±50% perturbation level, every $\mu_{m+\Delta P}^i$ is successively multiplied by 1.5, while the others remain unchanged. Multiplication by 0.5 for a −50% perturbation. A first observation was that due to nonlinearity, symmetrical perturbations did not result in symmetrical differences compared to the basal state (Fig. 1C and D). In particular, the squared error was greater with −50% perturbations than with +50% perturbations. Moreover, we observed that connectivity coefficients that were zero in the basal state, were no more zero when estimated with standard MRA (Fig. 1B–D). Departure from zero was computed: the perturbation was applied to each node successively, $r_{ij}$ equals $\sqrt{\sum (r_{ij}^0 - r_{ij})^2}$. In Fig. 2A, we see that it is nonsymmetric as expected from the results above. To limit errors in $r_{ij}$ estimates, one could try using smaller perturbations, but as already mentioned, in the presence of noise, this approach might be inappropriate. To illustrate this phenomenon, three levels of Gaussian additive noise $N(0, \sigma)$ were hence added to the $X_i$ with $\sigma = kX$, where $X$ was the average of all the theoretical concentrations at steady state ($X = 84$ for the example 6-node network), and $k$ was set at 0.1%, 0.5%, or 1%. Note that with $k = 1\%$, a substantial noise is added to low signals ($k = 1\%$ implies $\sigma = 0.84$). The error in the presence of noise (Fig. 2B) behaved in a reverse fashion compared to what we observed without noise. This is because the noise error was larger than the nonlinearity error. With more noise, it is indeed necessary to increase perturbation intensities to obtain a sufficient signal-to-noise ratio, but this also drives us away from the exact solution in case the system is substantially nonlinear with respect to $P$ (Equation (4)). In what follows, we used ±50% perturbations as a reasonable compromise.

### 3.2 First comparison of methods
MRA assesses network edges quantitatively, but measurement noise usually generates nonzero values for all the connectivity coefficients. Figure 3A and B illustrate this phenomenon for 6-node kinase network introduced above. For standard MRA, we estimated 95% CIs applying a bootstrap with 100 repetitions. A connectivity matrix $r$ was obtained for each repetition of the bootstrap and CI boundaries were simply obtained as the 2.5th and 97.5th percentiles. We observed an already known relationship (Thomaseth et al. 2018): CI sizes increase with higher noise in the data, independent of the number of replicates. It is important to note that such CIs represent the consequence of the total variance (noise + nonlinearity), and by applying the standard MRA approach we are limited to such variance estimations a posteriori, e.g. via simulations such as the bootstrap. The independence on the number of replicates is an obvious limitation.

On the contrary, solving MRA with multilinear regression methods enables estimating the residual variance, i.e. the variance caused by the nonlinear contributions. Residual variance-based CIs is common practice when applying linear models to data. In Fig. 3A and B, we show such CIs for 3 and 5 replicates and noise levels at 0.1% and 0.5% of the average concentrations $X$ for LSE regression. Residual variance-based CIs were indeed strongly reduced compared to standard MRA total variance-based CIs, and more replicates yielded smaller CIs. In a previous study limited to 3-module networks (Thomaseth et al. 2018), the idea of applying LSE has been already envisioned. Nonetheless, LSE was applied to each replicate separately thus leading to poor performance. In one case, the authors linked dual perturbations and reported encouraging performance, but they did not follow-up on this observation.

### 3.3 Determining edges existence in the 6-node network
In some applications, it is only necessary to infer the existence of node interactions and the sign of the interaction (induction or inhibition). To consider such applications, we reduced the exact connectivity coefficient $r_{ij}$ to −1, 0, or 1 by simply taking sign($r_{ij}$). The output of MRA and MRA-regression methods were then also discretized to obtain values in {−1; 0; 1} (Section 2). Results are reported in Fig. 3C and Supplementary Table S1 (details of sensitivity and specificity computations are in Section 2 as well).
Already on this modest 6-node network, we see that MRA and MRA-regression performances depend on the noise level. Overall, the best sensitivity was achieved by STEP-Fo, LSE_CI, and LASSO (in this order), which outperformed MRA. TLR remained stable, regardless of the noise. Concerning specificity, the best results were obtained by MRA, STEP-Fo, and STEP-Bo. Upon noise increase, specificity was improved for MRA and LSE_CI because an increased noise inflated the CIs thus leading to the correct detection of every null edge (Supplementary Table S1).

3.4 Medium-size networks from DREAM 4 Challenge

The DREAM 4 Challenge released five 10-gene and five 100-gene networks of *Escherichia coli* meant to compare inference algorithm performance. Each gene was subjected to two independent perturbation series, knockdowns (KDs) and knockouts (KOs). All the gene expression levels $X_i$ were normalized and an unknown noise was added. The exact solutions were provided as binary edges, i.e. $r_{ij} \in \{0, 1\}$. The networks were sparse with 80% of null edges on average. DREAM 4 data regarding the network dynamics before reaching a steady state were ignored. We evaluated the performance of our MRA-regression methods on the existence or nonexistence of the inferred edges, thus requiring discretization of the output real-valued $r_{ij}$. In the case of MRA, it was not possible to combine KO and KD data in one analysis. We hence treated the two sets of perturbations separately. Regarding MRA output selection and discretization, we used $x = 20\%$ for 10-gene, and $x = 25\%$ for 100-gene networks (see Section 2). Figure 4A features the first 10-gene network compared with the result found by TLR, detailed results for each network are featured in Supplementary Table S2. Figure 4B reports the average performance over the five networks of each size in terms of sensitivity and specificity. To eliminate potential biases due to specific thresholds in the discretization, we also computed areas under the ROC curve (AUROC) featured in Fig. 4C.

On these larger networks, regressive approaches showed strong superiority over standard MRA (Fig. 4B and C). MRA-KD was close to random selection for 10- and 100-gene networks, whereas MRA-KO performed reasonably with 10-gene networks, but was also close to random with 100 genes (Supplementary Figs S1–S3). The simple regression method TLR delivered better performance, but was nonetheless sensitive to the network size. Regression methods adapted to sparse solutions (STEP-xx and LASSO) performed robustly and STEP-xx provided the best compromise overall. Comparing these methods with DREAM 4 Challenge ranking established in 2014, based on P-values concerning areas under the ROC curve (official criterion), we found that STEP-xx and TLR methods ranked 3rd for 10-gene networks. They respectively ranked 1st and 8th for 100-gene networks (STEP-Fo being the best of STEP-xx). See Supplementary Table S3 for detailed results and the exact score computation.

Naturally, these rankings do not take into account methods developed since DREAM 4. In particular, a variation of MRA called MLMSMRA (Klinger and Blüthgen 2018), using a likelihood estimation combined with a greedy hill-climbing model selection approach, ranked 3rd with 10-networks and KO perturbations, but 25th with KD. The authors did not report performance for 100-node networks. A Bayesian-modified version of MRA was proposed to better model sparse networks according to another ranking system defined by the authors themselves (Santra et al. 2013; Halasz et al. 2016; Klinger and Blüthgen 2018), i.e. achieving a performance comparable to MRA combined with STEP-xx.

3.5 Synthetic networks of sizes up to 1,000 nodes

We applied a realistic random biological network generator FRANK (Carre et al. 2017) to systematically explore performance on network sizes from 30 to 1,000 at two noise levels. We applied -50% (KD) and full KOs perturbations. We generated eight sets of networks with TF in (30, 60, 100, 200, 300, 500, 800, and 1000) and TA = 0. In FRANK terminology, TFs are genes regulating at least another gene (out-degree ≥ 0), whereas TAs harbor out-degrees = 0. We also generated seven sets of networks, with TF = TA in (30, 50, 100, 150, 250, 400, and 500). The number of nodes (TF + TA) varied from 30 to 1000. For each set, we generated five independent networks with two different perturbations (KO and KD) and two noise levels. This resulted in $5 \times (8 + 7) \times 2 = 150$ synthetic networks. FRANK sparsity parameter was set to have 85% of null edges. For each network and noise level, we computed the exact squared error $\sqrt{\sum (r_{ij} - r_{ij})^2}$ for standard MRA and two
regression-based methods that stood out in the DREAM 4 data, i.e. TLR and STEP-Fo. See Fig. 5 and Supplementary Table S4 for the results, which confirmed on this large collection of networks the significant improvement brought by the regression approach.

To enable comparison with the network inference algorithms ARACNE, CLR, and MRNET, we next asked about the performance just detecting the existence of edges, i.e. both the reference connectivity coefficient $r_{ij}$ and their estimates $\hat{r}_{ij}$ were discretized to $\{0, 1\}$ (Section 2). Since AUROC computation is very time consuming, we limited performance assessment with AUROC to MRA and TLR (Fig. 6), which again showed the advantage of the regressive approach.

In Fig. 7, we report the performance of all the methods by simply computing the distance of a single (sensitivity, $1 - $ specificity) point, obtained with default parameters, to the diagonal representing random selection. We clearly see that network inference methods do not perform well in comparison to MRA-based methods on perturbation data. We also see that standard MRA performs well with small noise but its performance drops dramatically as noise increases in line with previous results. The most robust methods are the simple TLR and STEP-Fo, but TLR outperforms STEP-Fo on this dataset, and it is much faster. LASSO performance is robust although seldom the best.

3.6 The use of a priori knowledge of the network topology
In practice, it is common to know about some interactions in a studied network. Regarding known edges, there is nothing particular
Figure 4  Dream 4 Challenge networks. (A) A 10-gene network topology (left) compared to TLR inference (right). (B) Average performance of the methods over the five 10- and five 100-gene networks. Se, sensitivity; Sp, specificity. (C) Average performance reported as area under the ROC curve (AUROC) and corresponding $P$-values over the five 10- and five 100-gene networks. *AUROC and $P$-values were computed using the library verification (function roc.area). For LSE_CI, it failed, and we estimated the AUROC by ourselves without $P$-value.

Figure 5  FRANK synthetic networks. (A) Squared error (SE) for MRA, TLR and STEP-Fo methods as a function of the number of nodes with additive Gaussian noise $N(0, \sigma)$ and $\sigma = \bar{X}$, $\bar{X}$ the average concentration, and $k = 10\%$, number of nodes $= TF$, TA = 0. (B) Same with TA = TF, number of nodes = TA + TF. (C) Log (1 + SE) as a function of the number of nodes, $k = 50\%$, number of nodes = TF, TA = 0. (D) Same with TA = TF, number of nodes = TA + TF. The ribbon around TLR and STEP-Fo curves features ± standard deviation of the error. For MRA, the error standard deviation was so large that it would encompass the whole plot. It was hence not represented. Since STEP-Fo was very compute-intensive, we limited its application to networks with TF $\leq$ 300 and TA = 0, or TF $\leq$ 100 and TA = TF.
to do since MRA formalism considers all the interactions as possible. On the other hand, by knowing impossible interactions, we can eliminate equations (see Section 2). This increases over determination of the linear system of the regression approach, which is expected to increase the accuracy of the estimates. For methods such as LASSO that require hyper-parameter adjustment through CV, this has the potential to improve the optimality of those hyper-parameters in addition.

In the 10- and 100-gene networks of the DREAM 4 Challenge, we assumed different percentages of known null edges. The impact of this knowledge on sensibility and specificity for three of the studied regression methods (LASSO, STEP-Fo, and TLR) is featured in Fig. 8. Indeed, performance improved as a function of the percentage of known null edges. This improvement was stronger for specificity. We observed high variability of the impact on sensibility for 10-gene networks, results were more stable for the larger 100-gene networks.

In comparison with the binary approach proposed here, a Bayesian implementation of MRA has been proposed (Santra et al. 2013) that took a known pathway topology as a prior and showed improved accuracy over pure MRA. This Bayesian formulation was less radical than imposing null edges since an absent edge of the pathway could be eventually inferred a posteriori, provided that sufficient data evidence existed. On the other hand, our regression-based solution provides accurate CI estimates that hence enable pruning network edges based on model accuracy. One can imagine following with a reduced model, where pruned edges would be removed as above, to obtain better estimates on the other edges. This would be different from a Bayesian formulation obviously, though it would provide comparable practical value.

In the case of multilinear regression, the integration of the knowledge of null edges is done naturally (overfitting the system by deleting columns). In the Bayesian approach (Santra et al. 2013), the integration of this knowledge is also very natural (defining the prior probability of the Boolean variables associated with the existence of edges). In MLMSMRA (Klinger and Blu¨thgen 2018), the integration of prior knowledge was not explicitly discussed. Nonetheless, because their algorithm fills the connectivity matrix iteratively, it should be easy to force certain coefficients to remain at zero.

4 Conclusion

We have introduced a new method to solve MRA equations through multilinear regression. This formulation brings a number of advantages over the classical approach by providing a natural way to model data variability across experimental replicates, or even multiple perturbations at certain or all the modules. Better estimates of MRA connectivity coefficient variability can also be exploited to identify absent edges in a biological network more accurately. Moreover, these advantages were obtained by remaining within the MRA formalism that provides an elegant, physical interpretation of connectivity coefficients compared to purely regressive approaches.

Figure 6 FRANK synthetic networks. (A) AUROC as a function of the number of nodes. Additive Gaussian noise $N(0, \sigma)$ with $\sigma = k \bar{x}$, $\bar{x}$ the average concentration, and $k = 50\%$. Number of nodes = TF, TA = 0. (B) Same with $TA = TF$, number of nodes $= TA + TF$. See Supplementary Table S5 for detailed results

Figure 7 FRANK synthetic networks. (A) Distance to the diagonal as a function of the number of nodes with additive Gaussian noise $N(0, \sigma)$ with $\sigma = k \bar{x}$, $\bar{x}$ the average concentration, and $k = 10\%$. Number of nodes = TF, TA = 0. (B) Same with $TA = TF$, number of nodes $= TA + TF$. (C) Distance to the diagonal as a function of the number of nodes, $k = 50\%$, number of nodes = TF, TA = 0. (D) Same with $TA = TF$, number of nodes $= TA + TF$. (E) Distance of a point on a ROC curve to the diagonal. See Supplementary Table S6 for detailed results.
While motivating the use of regression for MRA, we conducted an analysis of the relationship between data noise, perturbation intensity, and MRA result accuracy that has interest per se. In agreement with the local development in the Taylor series, MRA accuracy decreases with stronger perturbations as soon as the system is nonlinear. On the other hand, the presence of noise requires perturbations with a minimal intensity to obtain exploitable differences in the variables. Altogether, this means that a compromise must be found between noise levels and perturbation strengths. Depending on the biological system at hand, i.e. on the nonlinearities, this may lead to perturbations intensity no longer compatible with MRA linear approximation. In such a case, another modeling paradigm must be chosen. In the absence of strong nonlinearities, our work dramatically extended the domain of the application of MRA to much larger networks of sizes up to 1000. This is a 100-fold increase compared to MRA with standard linear algebra, which had difficulties going beyond 10-node networks in our experiments.

Finally, the proposed approach actually defines a family of MRA-derived methods with the multilinear regression algorithm as a free parameter. While LSE_CI for instance selects edges based on a direct exploitation of residual variance CIs, other algorithms perform model selection such as the simple although very efficient TLR, LASSO, or STEP-xx. Other regression methods that we have not tested might provide specific advantages depending on the dataset, or the specific interest or requirements of the researchers.

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Supplementary data
Supplementary data is available at Bioinformatics online.

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References
Andrec M, Kholodenko BN, Levy RM et al. Inference of signaling and gene regulatory networks by steady-state perturbation experiments: structure and accuracy. J Theor Biol 2005;232:427–41.
Carré C, Mas A, Krouk G et al. Reverse engineering highlights potential principles of large gene regulatory network design and learning. NIPJ Syst Biol Appl 2017;3:17.
Dorel M, Klinger B, Gross T et al. Modelling signalling networks from perturbation data. Bioinformatics 2018;34:4079–86.
Faith JJ, Hayete B, Thaden JT et al. Large-scale mapping and validation of Escherichia coli transcriptional regulation from a compendium of expression profiles. PLoS Biol 2007;5:e8.
Halasz M, Kholodenko BN, Kolch W et al. Integrating network reconstruction with mechanistic modeling to predict cancer therapies. Sci Signal 2016;9:ra114.
Hastie T, Tibshirani R, Friedman J. The Elements of Statistical Learning: Data Mining, Inference, and Prediction. New York, NY: Springer, 2009.
Huynh-Thu VA, Sanguinetti G. Gene regulatory network inference: an introductory survey. Methods Mol Biol 2019;1883:1–23.
Jimenez-Dominguez G, Ravel P, Jalaguier S et al. An R package for generic modular response analysis and its application to estrogen and retinoic acid receptor crosstalk. Sci Rep 2021;11:7272.
Kholodenko BN, Kiyatkin A, Bruggeman FJ et al. Untangling the wires: a strategy to trace functional interactions in signaling and gene networks. Proc Natl Acad Sci USA 2002;99:12841–6.
Klinger B, Bluthgen N. Reverse engineering gene regulatory networks by modular response analysis - a benchmark. Essays Biochem 2018;62:535–47.
Meyer PE, Kontos K, Lafitte F et al. Information-theoretic inference of large transcriptional regulatory networks. EURASIP J Bioinform Syst Biol 2007;2007:79879.
Santos SDM, Verveer PJ, Bastiaens PIH et al. Growth factor-induced MAPK network topology shapes Erk response determining PC-12 cell fate. Nat Cell Biol 2007;9:324–30.
Santra T, Kolch W, Kholodenko BN et al. Integrating Bayesian variable selection with modular response analysis to infer biochemical network topology. BMC Syst Biol 2013;7:57.
Santra T, Rukhlenko O, Zhernovkov V et al. Reconstructing static and dynamic models of signaling pathways using modular response analysis. Curr Opin Syst Biol 2018;9:11–21.
Thomaseth C, Fey D, Santra T et al. Impact of measurement noise, experimental design, and estimation methods on modular response analysis based network reconstruction. Sci Rep 2018;8:16217.