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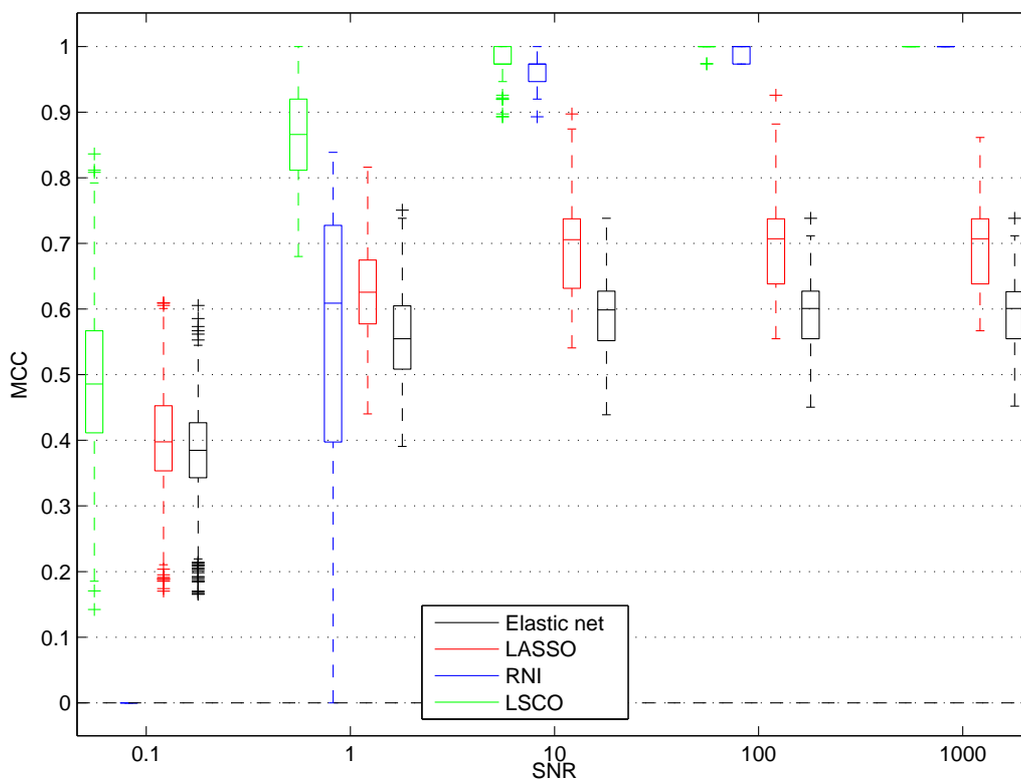
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L1 regularisation methods fail to infer the correct network even when the data is so informative that all existing links can be proven to exist.

# Avoiding pitfalls in $L_1$ -regularised inference of gene networks<sup>†</sup>

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Statistical regularisation methods such as LASSO and related  $L_1$  regularised regression methods are commonly used to construct models of gene regulatory networks. Although they theoretically can infer the correct network structure, they have been shown in practice to make errors, *i.e.* leave out existing links and include non-existing links.

We show that  $L_1$  regularisation methods typically produce a poor network model when the analysed data is ill-conditioned, *i.e.* the gene expression data matrix has a high condition number, even if it contains enough information for correct network inference. However, the correct structure of network models can be obtained for informative data, *i.e.* data with such a signal to noise ratio that existing links can be proven to exist, when these methods fail, by using least-squares regression and setting small parameters to zero, or by using robust network inference, a recent method taking the intersection of all non-rejectable models.

Since available experimental data sets are generally ill-conditioned, we recommend to check the condition number of the data matrix to avoid this pitfall of  $L_1$  regularised inference, and to also consider alternative methods.

## 1 Introduction

Gene regulatory network (GRN) inference, also known as reverse engineering or network reconstruction, is an essential endeavour in Systems biology. Several studies<sup>1–3</sup> state that mRNA transcriptional regulatory networks can be inferred based on gene expression data obtained from *in vivo* experiments in which all genes of interest are systematically perturbed and the resulting expression changes are measured. To be biologically realistic, the network needs to be relatively sparsely connected, in other words, only a fraction of all possible links exist. The LASSO method<sup>4</sup> and derivatives of it, all of which use  $L_1$ -regularisation to induce sparsity, achieve this and have become popular for GRN inference. Several other modelling techniques exist such as Bayesian<sup>5,6</sup>, information theoretic<sup>7,8</sup>, neural networks<sup>9,10</sup>, Boolean<sup>11,12</sup> and

dynamical systems<sup>1,13</sup>. Nonetheless, in this study we focus on  $L_1$ -regularisation methods, in particular LASSO, Elastic Net<sup>14</sup>, and Bolasso<sup>15</sup>, due to their widespread usage. We show that they fail to infer the correct network even when the data is informative enough for correct inference by other methods. We also test the methods on the *in vivo* data collected by Lorenz *et al.*<sup>2</sup> for inference of the *Snf1* network in *S. cerevisiae* and relate the result to our simulations on *in silico* data with known golden standard networks.

Theoretically, LASSO has been shown to be able to recover the correct network under certain conditions, such as the Strong Irrepresentable Condition (SIC) and Restricted Isometry Property (RIP)<sup>16–18</sup>. In a network inference context, these conditions concern the relation among observed vectors of expression changes. However, even results based on SIC only ensure that the LASSO estimator is sign consistent with a probability that goes to one as the number of samples goes to infinity. Some of the inferred links could thus not exist in reality, in particular for the low number of samples seen in biological data sets. In real applications, SIC is of little use because it cannot be calculated without knowing the true network. Even though performance of  $L_1$ -regularisation methods has been analysed rather extensively, we have not seen any article reporting that they fail for sufficiently informative data, which we show here.

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In a number of cases, when reverse engineering algorithms have been applied to biological networks, believed to have a well understood connectivity, networks with a different connectivity have been obtained. For instance, Lorenz *et al.*<sup>2</sup> reported a mere 62% sensitivity and 69% precision with 24% of the predicted regulatory interactions having the opposite sign in the model of the *Snf1* network in *S. cerevisiae*. Moreover, benchmarking studies, such as the Dialogue for Reverse Engineering Assessments and Methods (DREAM), have shown that GRN inference usually results in a large fraction of false positives, *i.e.* inferred links absent in the true network, and false negatives, *i.e.* missed links present in the true network<sup>19,20</sup>. This has in later years lead researchers to complement expression data with other data types, such as binding data, ChIP-seq, and *a priori* information<sup>21,22</sup>. Note that we here speak about addition of other data types to guide the inference method and not integration of other data types in the model. In the former case the degree of freedom of the model is kept fixed and the data is intended to constrain model parameters, while the degree of freedom in the later case is increased. Use of these, so called multi-data-type genomic datasets, makes it harder to assess the performance of inference methods compared to expression data alone. It is in particular harder to know to which degree a link is supported by expression data versus *a priori* information. Even if the complete topology of the network is provided, *e.g.* from ChIP binding data, the signs (activation/repression) of the links still need to be inferred. Addition of other data does not fix the method *per se*. We therefore think that awareness of the pitfall of  $L_1$ -regularisation methods that we report here is more essential than before.

A number of GRN inference benchmark studies<sup>23–25</sup> have been published, spanning a wide range of methods and data sets. In general, the conclusion is that although they tend to perform better than random, all inference methods produce models that are far from correct. The dependency on the nature of the data is strong as a method may do well in one benchmark but poorly in another one. Selection of the regularisation coefficient, which determines the sparsity of the estimate, is a major issue because it must be correct for the estimated network model to be correct<sup>26</sup>. Vinh *et al.*<sup>27</sup> detail the difficulties of benchmarking, especially on small networks, where sparsity cannot be achieved to any larger degree due to the network's small size. They show that methods for inference of GRNs do not construct any good networks with sufficient confidence and that the parameter settings of the algorithms are crucial to find a good estimate of the structure of the network. However, no method for optimising these crucial parameters is given. Jörnsten

*et al.*<sup>28</sup> show that the structural agreement between network models inferred for the same biological system using bootstrapping based on measurements obtained at two different platforms only is good for a narrow range of the regularisation coefficient. This makes it important to assess how the accuracy of different inference methods depends on data and system properties, which we here do for five methods.

Data sets generated *in vivo* for gold standard networks are rare for benchmark purposes due to a lack of knowledge about the interactions among the genes. An attempt has been made to create such a gold standard for benchmarking by recording an *in vivo* data set from a synthetically engineered five gene network in yeast, called IRMA<sup>3</sup>. Penfold and Wild<sup>24</sup> benchmarked time series algorithms in addition to steady-state algorithms and evaluated their performance on IRMA. They found that no methods could retrieve the designed structure of IRMA from the data. The IRMA network was perturbed by single gene over-expression to trigger the response of the network and the change in mRNA abundance was then measured when the system had reached steady-state, as well as a time series sampled either every 10 or 20 minutes for up to 5 hours. For single gene perturbations there is no guarantee that the gene space is sufficiently excited to give informative data, *i.e.* that a sufficient variation in the response of the genes over the experiments is achieved<sup>29</sup>. Another issue with gold standard networks is the definition of a link. The inference method and model formalism has to yield the same type of links as recorded in the gold standard in order for a comparison to be meaningful and fair. The five methods employed here infers so called influences, while gold standard networks typically contain links corresponding to physical binding between molecules<sup>23</sup>. Simulated data sets are thus still necessary for benchmarking due to the lack of “real” data sets that are informative enough for accurate GRN inference and differences in the definition of a link. It is thus not possible to exhaustively demonstrate the pitfalls of  $L_1$ -regularisation methods on real data, despite the multitude of data that exist. However, we have applied the studied inference methods to the *in vivo* data collected by Lorenz *et al.*<sup>2</sup>, compared the inferred networks to three networks that can be seen as gold standards, and relate the accuracies to the expected performance in our simulations based on the properties of the data.

In this study, we focus on analysing network and data properties that are important for the accuracy of GRN inference. In particular, the condition number of the network and response matrices, as well as the Signal to Noise Ratio (SNR), are examined. To this end, we generated a set of linear networks with essential properties simi-

lar to real biological GRNs. These were then used to generate both gene expression data sets that have properties similar to published *in vivo* data and data sets that are informative enough for inference of the correct network. This was done to mimic real data sets, while varying the properties and utilising the advantage of knowing the true network. We restrict ourselves to linear models, because it is sufficient to demonstrate the presented pitfall of  $L_1$ -regularisation methods. Considering that the class of linear models is a subset of the class of nonlinear models, awareness of this pitfall is essential also when inferring a nonlinear model. By identifying easily testable conditions that need to be satisfied for successful GRN inference, we provide guidelines useful for avoiding pitfalls that can cause poor network models.

## 2 Problem description

In this paper we make the common assumption that the GRN can be described by a linear dynamical systems model<sup>1,30,31</sup>

$$\begin{aligned} \dot{x}_i(t) &= \sum_{j=1}^N a_{ij}x_j(t) + p_i(t) - f_i(t) \\ y_i(t) &= x_i(t) + e_i(t). \end{aligned} \quad (1)$$

In biological terms, the state vector  $\mathbf{x}(t) = [x_1(t), x_2(t), \dots, x_N(t)]^T$  represents actual mRNA expression changes relative to the initial state of the system, the perturbation vector  $\mathbf{p}(t) = [p_1(t), p_2(t), \dots, p_N(t)]^T$  represents the applied perturbation, which may be corrupted by the noise  $\mathbf{f}(t)$ . The perturbations could be *e.g.* gene knock-downs using siRNA or gene over-expressions using a plasmid with an extra copy of the gene. The response vector  $\mathbf{y}(t) = [y_1(t), y_2(t), \dots, y_N(t)]^T$  represents the measured expression changes that differ from the true expression changes by the noise  $\mathbf{e}(t)$ . The parameters  $a_{ij}$  of the interaction matrix describe the influence of an expression change of gene  $j$  on gene  $i$ . A positive value represents an activation, while a negative value represents an inhibition. The relative strength of the interaction is given by the value of the  $a_{ij}$  parameter. We make the common assumption that only steady-state data is recorded, which simplifies our data model (1) to

$$\mathbf{Y} = -\mathbf{A}^{-1}\mathbf{P} + \mathbf{A}^{-1}\mathbf{F} + \mathbf{E} \quad (2)$$

when the set of experiments are considered. Here  $\mathbf{Y}$  is the observed steady-state response matrix after applying the perturbations  $\mathbf{P}$  and  $\mathbf{A}$  is the interaction matrix *i.e.* network.

By taking the transpose of the variables and “true” network model, and introducing the notation used for regressors  $\check{\Phi} \triangleq [\phi_1, \dots, \phi_j, \dots, \phi_N] = \mathbf{Y}^T$ , regressands

$\Xi \triangleq [\xi_1, \dots, \xi_i, \dots, \xi_N] = -\mathbf{P}^T$ , regressor errors  $\Upsilon \triangleq [\nu_1, \dots, \nu_j, \dots, \nu_N] = \mathbf{E}^T$ , and regressand errors  $\Pi \triangleq [\epsilon_1, \dots, \epsilon_i, \dots, \epsilon_N] = -\mathbf{F}^T$ , we obtain the matrix form of the standard linear data model used in errors-in-variables regression problems

$$\check{\Phi} = \check{\Phi} + \Upsilon, \quad \Xi = \check{\Xi} + \Pi \quad (3a)$$

$$\check{\Phi}\check{\mathbf{A}}^T = \check{\Xi} \quad \check{\Phi}, \check{\Xi} \in \mathbb{R}^{M \times N}. \quad (3b)$$

Here  $M$  is the number of experiments/samples, *i.e.* data points, and  $N$  is the number of states/nodes.

## 3 Materials and methods

### 3.1 Network inference algorithms

Least Absolute Shrinkage and Selection Operator (LASSO) penalises models with small nonzero parameters by introducing a  $L_1$  penalty term in the objective function which equals the sum of the absolute values of the parameters<sup>4</sup>

$$\hat{\mathbf{A}}_{\text{reg}}(\check{\zeta}) = \arg \min_{\mathbf{A}} \|\mathbf{A}\mathbf{Y} + \mathbf{P}\|_{L_2}^2 + \check{\zeta} \|\mathbf{A}\|_{L_1}. \quad (4)$$

The effect of the introduced  $L_1$  regularisation term depends on the regularisation parameter  $\zeta$ . If it is set to zero then the ordinary least squares estimate is obtained, while a network model with no links is obtained when it goes to infinity. The regularisation term will trade the models predictive performance on the fitted data for a reduction of the number of descriptive model parameters.

The Elastic net<sup>14</sup> is a method based on LASSO which combines the  $L_1$  penalty from LASSO and the  $L_2$  penalty from ridge regression. The influence of the penalties are then weighted by a parameter  $\alpha$  such that,

$$\hat{\mathbf{A}}_{\text{reg}}(\check{\zeta}) = \arg \min_{\mathbf{A}} C + \check{\zeta} (\alpha \|\mathbf{A}\|_{L_1} + (1 - \alpha) \|\mathbf{A}\|_{L_2}^2), \quad (5)$$

where  $C = \|\mathbf{A}\mathbf{Y} + \mathbf{P}\|_{L_2}^2$ .

Bolasso<sup>15</sup> is a bootstrap approach to LASSO inference, where the statistical properties of bootstrapping are combined with the LASSO, see algorithm 1. We use a constant number of bootstraps,  $n_{BS} = 100$ , for each data set, as the statistical confidence should increase with  $n_{BS}$ . This is well above the minimum number of bootstraps needed<sup>15</sup>,  $\sqrt{N}$ , with  $N$  being the number of variables. We extend the bootstrap algorithm by requiring that the bootstrapped data set has the same rank as the original data. In practice this means putting a rank requirement on the  $\mathbf{P}$  matrix so that it has full row rank. This improves the performance, because it ensures that all genes are perturbed in at least one experiment, which is a necessary condition for correct inference<sup>32</sup>. Bolasso was not

**Algorithm 1** Plain bootstrap LASSO algorithm.  $\mathbf{B}$  is the inferred model and  $\mathbf{A}$  the logical intersection of inferred models.  $a_{ij}$  is a link from  $j$  to  $i$ .  $n_{BS}$  is the number of bootstraps.

---

```

procedure BOOTSTRAP LASSO(data,  $n_{BS}$ )
   $a_{ij} = 1 \forall i$  and  $j \in \mathbf{A}$ 
  for  $1:n_{BS}$  do
    dataBS = DRAW WITH REPLACEMENT(data)
     $\mathbf{B} = \text{LASSO}(\text{bdata}_{BS}, \zeta)$ 
     $\mathbf{A} = \mathbf{A} \wedge \text{LOGICAL}(\mathbf{B})$ 
  end for
   $\mathbf{A} = \{a_{ij} \in \mathbf{A}\}$ 
end procedure
function DRAW WITH REPLACEMENT(data)
  Draw samples with replacement
  s.t.  $|\text{data}_{BS}| = |\text{data}|$ 
  return dataBS
end function

```

---

applied to the 10 gene data sets because the data matrix becomes rank deficient if a sample that is left out during the bootstrap procedure contains the only perturbation of a gene. This is often the case in the 10 gene data, and the consequence is that links can not achieve 100% bootstrap support if they can only be inferred when that unique experiment is sampled. For the same reason, Blasso was not applied to the data by Lorenz *et al.*<sup>2</sup> as it only consists of one set of single gene perturbations, leading to a rank deficient data matrix as soon as one of the experiments is excluded during the bootstrap procedure.

Least-Squares Cut-Off (LSCO) is a simple inference algorithm based on ordinary least squares (OLS) followed by the removal of all weak links, *i.e.* small nonzero parameters,

$$\hat{a}_{ij} \triangleq \begin{cases} a_{ij}^{ols} & \text{if } |a_{ij}^{ols}| \geq \tilde{\zeta} \\ 0 & \text{otherwise} \end{cases} \quad \text{with } \mathbf{A}_{ols} \triangleq -\mathbf{P}\mathbf{Y}^\dagger. \quad (6)$$

The cutoff is used like a sparsity parameter and is varied over a range; for each data set the value producing the network with structure closest to the true network was picked<sup>26</sup>.

Robust Network Inference (RNI) is achieved by implicitly checking all network models that cannot be rejected based on the assumed data model and desired significance level and only including the links that are present in all of these models<sup>32</sup>. This gives the intersection of all non-rejectable models. In practice, the network model is obtained by calculating Nordling's confidence score and only including links with a value above one. Nordling's confidence score for the existence of the link  $a_{ij}$  is defined

as

$$\gamma(a_{ij}) \triangleq \sigma_N(\Psi(\chi)), \quad (7a)$$

with each element

$$\psi_{kl}(\chi) \triangleq \frac{\psi_{kl}}{\sqrt{\chi^{-2}(\alpha, NM)\lambda_{kl}}} \quad (7b)$$

$$\text{and } \Psi \triangleq [\phi_1, \dots, \phi_{j-1}, \phi_{j+1}, \dots, \phi_N, \xi_i], \quad (7c)$$

assuming that the data has been generated by the data model (3), with  $v_j$  and  $\epsilon_i$  drawn from a normal distribution with zero mean and a diagonal covariance matrix<sup>32</sup>. Here  $\sigma_N$  denotes the  $N$ th singular value, and  $\chi^{-2}(\alpha, NM)$  the inverse of the chi-square cumulative distribution with  $NM$  degrees of freedom, such that  $\text{P}[\chi^2(NM) > \chi^{-2}(\alpha, NM)] = \alpha$ <sup>33</sup>. A confidence score above one implies that the link can be proven to exist at the desired significance level  $\alpha$ , in this article set to 0.01. RNI obtains a network model that under the assumptions above only contains true positives<sup>32</sup>. False positives are thus avoided at the expense of accepting false negatives. RNI was done using code provided by Nordron AB ([www.nordron.com](http://www.nordron.com)), which owns all rights.

### 3.2 Networks

To assess the performance of the inference methods we generated a number of networks by varying model properties that have been considered important in the literature<sup>29,34–36</sup>. The sparsity of the networks was set to 0.25 for  $N = 10$  based on reported sparsities. For instance, the data on the ten gene network of the Snf1 signalling pathway in *S. cerevisiae*<sup>2</sup> can be explained well with networks having a sparsity in the range 0.22 to 0.28 and 29 transcriptional regulatory influences have been reported for it in the literature. For  $N = 45$  we generated networks with a sparsity around 0.07. Sparsity is defined as the fraction of links present in the network, denoted  $L$ , relative to the total number of possible links,  $N^2$ , *i.e.*  $s \triangleq \frac{L}{N^2}$ . The interampattness degree for a linear system is defined as the condition number of the system matrix  $\mathbf{G} = \mathbf{A}^{-1}$ <sup>29</sup>. It is thus  $\kappa(\mathbf{G}) \triangleq \kappa(\mathbf{A}) \triangleq \frac{\sigma_1(\mathbf{A})}{\sigma_N(\mathbf{A})}$ , where  $\sigma_1(\mathbf{A})$  and  $\sigma_N(\mathbf{A})$  are the largest and smallest singular values of the network matrix  $\mathbf{A}$ , respectively, for each network. We picked a small value between  $\kappa \in [0.5, 1] \cdot N$ , and a large value  $\kappa \in [9, 11] \cdot N$ , with 10 networks for each level. The latter is within the range reported for real networks based on data on a ten gene network of the Snf1 signalling pathway in *S. cerevisiae* 253, and a nine gene subnetwork of the SOS pathway in *E. coli* 54<sup>1,2,29</sup>. We generated the networks randomly, while making sure the networks have full rank, and weighted the model parameters to ensure stability<sup>37</sup>, and that we achieved the desired  $\kappa(\mathbf{A})$ .

Table 1 and Table 2 gives an overview of the  $N = 10$  and  $N = 45$  network properties respectively. For a complete list of the networks and properties see Table S.3 and Table S.4.

**Table 1** Network properties, for  $N = 10$  networks

| Network properties                           | low $\kappa(\mathbf{A})$ | high $\kappa(\mathbf{A})$ |
|--|--------------------------|---------------------------|
| # genes, $N$                                 | 10                       | 10                        |
| # networks                                   | 10                       | 10                        |
| structure                                    | random                   | random                    |
| interampattness degree, $\kappa(\mathbf{A})$ | 6.9 – 10                 | 91.6 – 108                |
| sparsity                                     | 0.25                     | 0.25                      |

**Table 2** Network properties, for  $N = 45$  networks

| Network properties                           | low $\kappa(\mathbf{A})$ | high $\kappa(\mathbf{A})$ |
|--|--------------------------|---------------------------|
| # genes, $N$                                 | 45                       | 45                        |
| # networks                                   | 10                       | 10                        |
| structure                                    | random                   | random                    |
| interampattness degree, $\kappa(\mathbf{A})$ | 25.4 – 41.3              | 411.5 – 492.8             |
| sparsity                                     | $\approx 0.07$           | $\approx 0.07$            |

### 3.3 Data sets

Data sets were created according to the linear dynamical model in (2). Given the true network  $\check{\mathbf{A}}$  we calculate an initial  $\mathbf{P}$  generated with the given perturbation design. Our noiseless expression data is then  $\check{\mathbf{Y}} = -\check{\mathbf{A}}^{-1}\mathbf{P}$ . We included  $2N$  samples for  $N = 10$  and  $4N$  samples for  $N = 45$ , because published data sets typically contain one to three replicates of  $N$  experiments<sup>1–3</sup>.

We followed three different perturbation approaches two for  $N = 10$ : Naive Random Double Perturbation (NRDP) and Sparse Balanced Excitation Design (SBED), and one for  $N = 45$ : triple Single Sets and a Single Double set (SSSD).

NRDP was constructed by perturbing two randomly chosen genes for each sample while making sure that  $\mathbf{P}$  had full rank and that each gene was perturbed at least once. By perturbing genes more than once we make sure that each sample has some dependency on the remaining data set, a requirement for using the sample in leave one out cross-optimisation of  $\zeta$ <sup>26</sup>. This design yields data sets where the condition number of  $\mathbf{Y}$  is close to the interampattness degree of the network. We therefore generate data sets with similar conditions to those reported in the literature, 5, 154, and 215, respectively in Gardner *et al.*<sup>1</sup>, Alter *et al.*<sup>34</sup>, and Lorenz *et al.*<sup>2</sup>.

The objective of the SBED is to excite all directions of the gene space uniformly, *i.e.* spread out the response

equally in the gene space, and obtain a well conditioned  $\mathbf{Y}$  matrix<sup>29</sup>. We do this approximately by minimising  $\kappa(\mathbf{Y})$  and the number of perturbed genes. To achieve uniform excitation is simpler for a dense perturbation matrix  $\mathbf{P}$  as the different signal directions in  $\mathbf{Y}$  can be more easily tuned. However as the possibility to perturb a majority of the genes at once is unrealistic, we keep  $\mathbf{P}$  as sparse as possible, *i.e.* we do a trade-off between a sparse perturbation design and uniform excitation in all directions of the gene space.

The SSSD perturbation design is constructed by using triple replicates where a single gene is perturbed for each sample with one extra set of double perturbation where two random genes are perturbed for each sample. This setup simulates a plausible experimental design approach that naively tries to maximise the information in the data set while utilising the fact that there need to be a dependence between samples to do some form of cross validation.

Table 3 and Table 4 shows an overview of data set properties. For a complete list of the data sets and properties see Table S.1, Table S.2, and Table S.5.

**Table 3** Data set properties

| Data set property                      | SBED      | NRDP        |
|--|-----------|-------------|
| perturbation design                    | SBED      | NRDP        |
| samples, $M$                           | $2N$      | $2N$        |
| # data sets                            | 20        | 20          |
| condition number, $\kappa(\mathbf{Y})$ | 1.3 – 2.0 | 9.5 – 181.3 |
| max # perturbations per sample         | 2 – 6     | 2           |
| min # perturbations per sample         | 1 – 3     | 2           |

**Table 4** Data set properties

| Data set property                      | SSSD        | SSSD           |
|--|-------------|----------------|
| perturbation design                    | SSSD        | SSSD           |
| samples, $M$                           | $4N$        | $4N$           |
| # data sets                            | 10          | 10             |
| condition number, $\kappa(\mathbf{Y})$ | 25.7 – 41.3 | 412.8 – 504.51 |
| max # perturbations per sample         | 2           | 2              |
| min # perturbations per sample         | 1           | 1              |

We applied noise to each data set with a variance  $\lambda$  selected to give the desired Signal to Noise Ratio (SNR)

$$\text{SNR} \triangleq \frac{\sigma_N(\check{\Phi})}{\sqrt{\chi^{-2}(\alpha, NM)\lambda}}, \quad (8)$$

where  $\sigma_N(\check{\Phi})$  is the  $N$ th singular value of  $\check{\Phi}$ , and  $\chi^{-2}(\alpha, NM)$  is the inverse of the chi-square cumulative distribution function as explained above. We generated

100 different noise realisations to do Monte Carlo simulations, each from a normal distribution with zero mean and variance  $\lambda$  using the `randn` function in Matlab version R2012a ([www.mathworks.com](http://www.mathworks.com)). For each data set, the variance of each realisation was then scaled based on (8) to achieve the desired SNR. For the data sets we used the significance level  $\alpha = 0.01$ . By covering the whole range of SNRs from completely uninformative to informative enough, we include the levels seen in real data.

### 3.4 Performance evaluation

We assessed the accuracy of the estimated networks using the Matthew Correlation Coefficient (MCC)<sup>38</sup>. MCC accounts for both true positives (TP), false positives (FP), true negatives (TN), and false negatives (FN), providing one number in the range  $[-1, 1]$  that captures the structural similarity between two networks containing the same labelled nodes. To use it one needs a golden standard that is taken as the true network that all the estimates are compared to.

The Fraction of Provably Existing Links (FPEL) is the fraction of links existing in the true network that can be proven to exist based on the observed data. They are proven to exist by rejecting all alternative network models lacking these links at a desired significance level based on the observed data and true data model, *i.e.* when the considered set of network models contains the true network and the measurement noise is described by the error model that was used to generate it. FPEL is calculated as the number of links with Nordling's confidence score (7) above one divided by the number of links in the true network<sup>32</sup>. It is the sensitivity of RNI. If all existing links can be proven to exist, *i.e.* FPEL = 1, then the data set is said to be informative enough for network inference. Note that FPEL and MCC are not directly comparable, since only the later accounts for FP and TN. MCC is relative to the number of possible links  $N^2$ , while FPEL is relative to the number of links present in the true network  $L$ . Nonetheless, MCC = 1 corresponds to FPEL = 1 for RNI. Only measurement data and an error model is needed to calculate the number of provably existing links, implying that it can be used for validation even when no golden standard or true network exists.

### 3.5 Analysis of the irrepresentable condition

The network model in (3b) can for each row  $i$  of the interaction matrix  $\mathbf{A}$  be expressed as  $\check{\Phi}\check{\theta}_i = \check{\xi}_i$ , yielding a sparse estimation problem for each row. By introducing  $\Phi_{0_i}$  and  $\Phi_{0_i^c}$  that contain regressors corresponding to the zero and nonzero elements of  $\check{\theta}_i$ , respectively, and  $\beta_i$

containing the nonzero elements of  $\check{\theta}_i$ , the Common part of the Irrepresentable Conditions (CIC), used by Zhao and Yu<sup>16</sup> in theorems ensuring sign consistency of the LASSO estimator, can be expressed as

$$\tilde{\mu}_i \triangleq \left| \Phi_{0_i}^T \Phi_{0_i^c} \left( \Phi_{0_i^c}^T \Phi_{0_i^c} \right)^{-1} \text{sign}(\beta_i) \right|. \quad (9)$$

If all elements of  $\tilde{\mu}_i$  are smaller than 1, then the Weak Irrepresentable Condition (WIC) is fulfilled and if all elements are smaller than 1 minus a positive constant  $\eta$ , then the Strong Irrepresentable Condition (SIC) is fulfilled<sup>16</sup>. The latter is used to show that LASSO is strongly sign consistent and the former that it is general sign consistent; both imply that the probability that all elements in the LASSO estimate of  $\theta_i$  have the correct sign goes to one when the number of samples  $M$  goes to infinity. A few additional technical conditions are required in the theorems, but it is logical to expect a high accuracy of the network estimate produced by LASSO when

$$\mu \triangleq \max_i \max_j \tilde{\mu}_i < 1 \quad (10)$$

If all columns in  $\Phi_{0_i}$  are orthogonal to all columns in  $\Phi_{0_i^c}$ , then  $\tilde{\mu}_i = 0$  and SIC is fulfilled. Assume for a moment that  $\Phi_{0_i} = \phi_1$  and  $\Phi_{0_i^c} = \phi_2$ , then  $\tilde{\mu}_i = \left| \phi_1^T \phi_2 / \|\phi_2\|^2 \right|$ . Now if  $\phi_1 = \alpha \phi_2$ , *i.e.*  $\phi_1$  is parallel to  $\phi_2$ , then  $\tilde{\mu}_i = |\alpha|$  is greater or equal to one unless  $\alpha$  is smaller than one, *i.e.* unless  $\phi_1$  is shorter than  $\phi_2$ . Hence the projection of any regressor corresponding to a zero element that is not orthogonal to the regressors corresponding to a nonzero element onto them must always be shorter than all of them to fulfill SIC. This would always hold if all regressors corresponding to a zero were shorter than all regressors corresponding to a nonzero element. This makes it interesting to calculate the minimum ratio between the shortest regressor corresponding to a nonzero element and the longest corresponding to a zero over all rows

$$r_{\min} \triangleq \min_i \left( \frac{\min_{\phi_k \in \Phi_{0_i^c}} \|\phi_k\|}{\max_{\phi_l \in \Phi_{0_i}} \|\phi_l\|} \right). \quad (11)$$

## 4 Results and discussion

We first present a comparison of the accuracy of the network models yielded by LASSO, Elastic Net, LSCO, and RNI as a function of the SNR for two different groups of data sets from 10 gene networks. One set generated by SBED in which the condition number of the response matrix,  $\mathbf{Y}$ , is low and another generated by NRDP in which it is high. Similarly, we also compare the accuracy of the

network models yielded by LASSO, Elastic Net, Bolasso, LSCO, and RNI for data sets from 45 gene networks. We used the network inference methods LASSO, Elastic Net, Bolasso as representatives of commonly used algorithms. In addition, we used LSCO and the recently proposed method RNI, which, under the assumptions used to generate our data sets, finds all links that can be proven to exist. We then use the irrepresentable conditions to analyse why and when LASSO fails based on these two groups of data sets. Finally, using the *in vivo* data collected by Lorenz *et al.*<sup>2</sup>, we demonstrate how our simulated results can be used to estimate the performance of the methods when they are applied to biological data.

#### 4.1 Vulnerability analysis of GRN inference methods

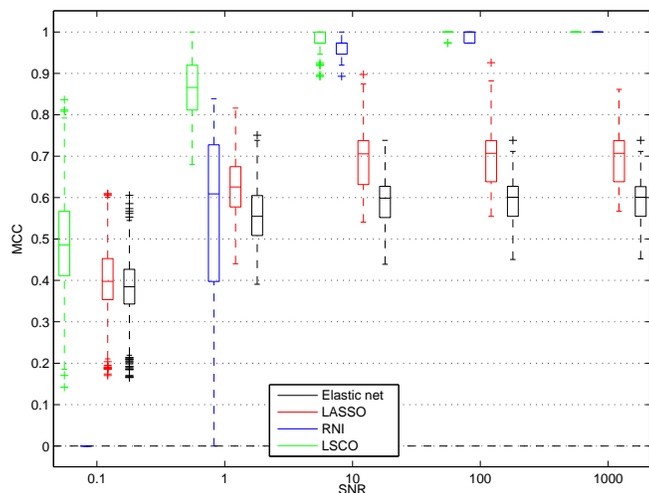
The most striking result on the data set with high response matrix condition number is that all the  $L_1$  regularisation methods fail to recover the true network model even when the SNR is so high that the data is informative enough for network inference and all existing links can be proven to exist (Figure 1 and 3). This unexpected failure of  $L_1$  regularisation constitutes an important pitfall in network inference, since many inference methods use  $L_1$  penalties and gene expression data sets often have a high condition number<sup>29</sup>. Actually the condition numbers of the 20 response matrices with  $N = 10$  are in the range 9 to 181, which is modest compared to the condition number of recorded response matrices used for inference of GRNs, *e.g.* the ten gene network of the *Snf1* signalling pathway in *S. cerevisiae* (215)<sup>2</sup> and the nine gene sub-network of the *SOS* pathway in *E. coli* (154)<sup>1</sup>. Even when the data is so informative that all existing links can be proven to exist using RNI, LASSO in the best case only obtained an MCC of 0.84, while LSCO in all cases for  $N = 10$  recovers the true network. In the  $N = 45$  case Bolasso outperforms LASSO and Elastic Net but fails to recover all links correctly even for the data sets that are informative enough. It is worth noting that Bolasso requires approximately 100 times more computations than LASSO. In LSCO the sum of squared residuals is minimised before any weak link is removed so it will provide good estimates for informative data. We therefore recommend all users of  $L_1$  regularisation to check the condition number of the response matrix in order to avoid this pitfall. If it is high, then LSCO and RNI can yield better network estimates.

It is important to note that in each case, for each noise realisation, we selected the  $\zeta$  value that yielded the LASSO estimate that was closest to the true network, *i.e.* highest MCC for the 100 noise realisations for each noise level, and similarly for Elastic Net, Bolasso, and LSCO.

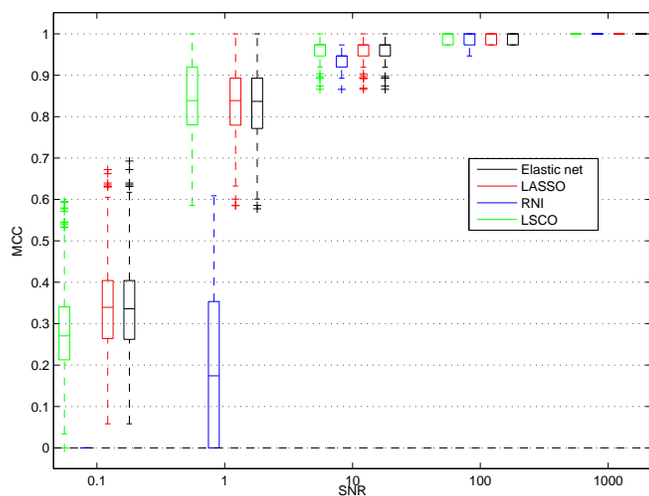
The former was done to avoid the influence of the rule used to select the regularisation coefficient  $\zeta$ , which typically has a strong influence on the accuracy of the network estimate and is difficult to select correctly<sup>26</sup>. Our network estimates are thus in general unrealistically accurate and require knowledge of the true network which is only available for simulated data, yet they are still far from correct. The latter was done to decrease the impact of random effects of the noise realisations in favour of data properties by doing Monte Carlo simulations. For this reason we also used the same 100 noise realisations for all data sets and all SNRs. We varied the network and data properties within ranges deemed reasonable and relevant for network inference based on previous works. In the literature a single gene is typically perturbed in each experiment, but we here used NRDP, *i.e.* perturbed two genes in each experiment, selected at random while ensuring that each gene is perturbed and that the perturbations constitute a linearly independent set. We also analysed data sets generated by the typical single gene design and observed the same failure of LASSO (data not shown). A total of  $2N$  ( $N = 10$ ) or  $4N$  ( $N = 45$ ) simulated perturbation experiments were used in all data sets, which is comparable to the  $3N$  experiments performed *in vivo* by Lorenz *et al.*<sup>2</sup> and Gardner *et al.*<sup>1</sup>, respectively.

For the 20 SBED data sets with a response matrix having a low condition number, LASSO, Elastic Net, and LSCO performed equally well and recovered the true network in all cases when the data sets were informative enough for network inference, see Figure 2. The SBED was in these cases used to balance the excitation of all directions in the space spanned by the 10 genes of the network, so that all singular values of the response matrix were of similar magnitude, while perturbing as few genes as possible in each experiment. These response matrices therefore have condition numbers in the range 1.3 to 2.0, which we have not yet seen for any published gene expression data set. It is worth remembering that we selected the optimal  $\zeta$  value for each data set and noise realisation for both LASSO, Elastic Net, and LSCO, so the performance is in general unrealistically good. For SNR 0.1, a weak indication of LASSO and Elastic Net outperforming LSCO is present but we cannot say that one method in practice is preferable over the other because the accuracy is sensitive to the selection of the value of the regularisation coefficient  $\zeta$ <sup>26</sup>. The same networks and noise realisations as described above were used.

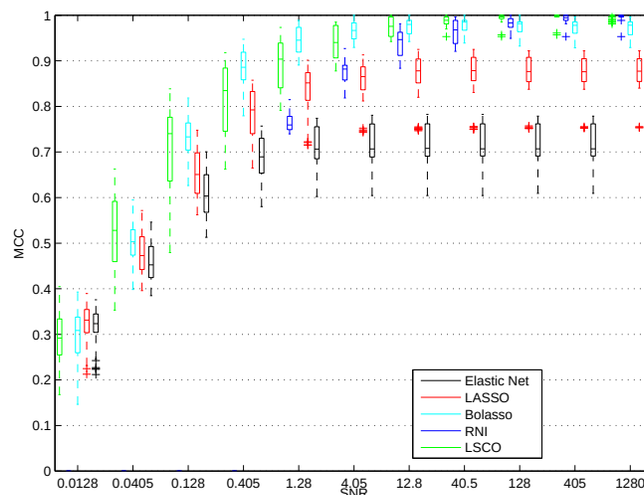
For the 10 SSSD data sets with a response matrix having a low condition number Bolasso and LSCO performed equally well and recovered the true network in all cases when the data sets were informative enough for network



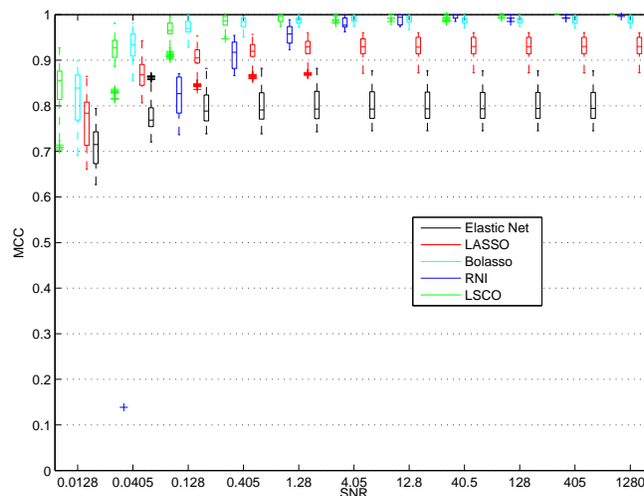
**Fig. 1** GRN inference accuracy versus signal to noise ratio using LASSO, Elastic Net, LSCO, and RNI on NRDP data sets with  $N = 10$  and high condition number  $\kappa(\mathbf{Y})$ . LASSO and Elastic Net fail even when all existing links can be proven to exist, corresponding to  $MCC = 1$  for RNI. Boxes are grouped according to five SNR values. Box edges signifies  $q_1 = 25$ th and  $q_3 = 75$ th percentile, whiskers encapsulate the most extreme data points not considered outliers. Outliers are considered points which are  $> q_3 + w(q_3 - q_1)$  or  $< q_1 - w(q_3 - q_1)$  where  $w = 1.5$  and marked with +.



**Fig. 2** GRN inference accuracy versus signal to noise ratio using LASSO, Elastic Net, LSCO, and RNI on SBED data sets with  $N = 10$  and low condition number  $\kappa(\mathbf{Y})$ . For an SNR of 10 LASSO, Elastic Net, and LSCO can infer the true network structure for some of the data sets even though all existing links cannot be proven to exist (RNI has a  $MCC < 1$ ). For an SNR  $> 10$  the median of all methods inference accuracy is approaching 1 and is above 90% for all data sets. For a description of the plot see Figure 1.



**Fig. 3** GRN inference accuracy versus signal to noise ratio for LASSO, Elastic Net, Bolasso, LSCO, and RNI on SSSD data sets with  $N=45$  and high condition number  $\kappa(\mathbf{Y})$ . All  $L_1$  regularised methods fail even when all existing links can be proven to exist, corresponding to  $MCC = 1$  for RNI. For a description of the plot see Figure 1.



**Fig. 4** GRN inference accuracy versus signal to noise ratio for LASSO, Elastic Net, Bolasso, LSCO, and RNI on SSSD data sets with  $N=45$  and low condition number  $\kappa(\mathbf{Y})$ . For a description of the plot see Figure 1.

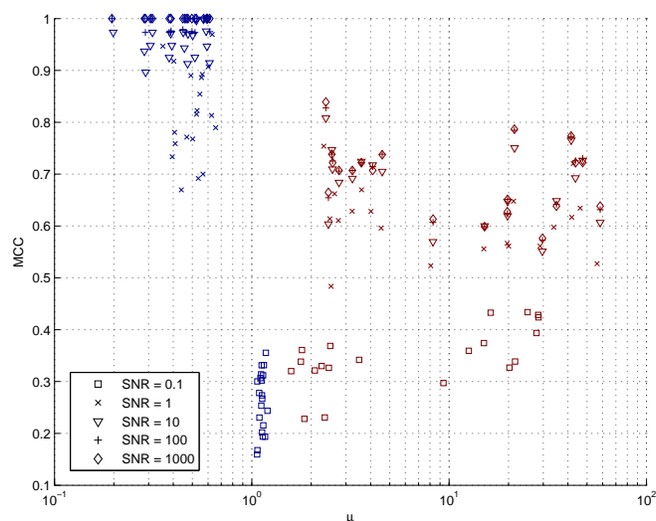
inference, see Figure 4, while LASSO and Elastic Net performed worse. This is probably due to the condition number of the response matrix being significantly larger than for the 10 gene case in Figure 2. It is now in the range 26 to 41. We also observe that Elastic Net performs worse than all other methods for the 45 gene case, but have not investigated why. The same networks and noise realisations as described above were used.

For low SNRs, RNI seems partly outperformed by all other methods because of a large number of false negatives, which are a consequence of ensuring that only true positives are included in the network model under the mild assumptions that are fulfilled here, and partly because the optimal regularisation coefficient  $\zeta$  is selected based on knowledge of the true network. RNI partly performs better than LASSO and Elastic Net from SNR 1 and better than Bolasso from SNR 100 for the data sets with a higher condition number because the SNR is defined based on the weakest singular value and the total excitation hence in general is higher. RNI is mainly included in this study because it gives FPEL and thereby can be seen as a lower bound on the performance that should be required from every other inference method. No other inference method that we are aware of can be used to prove that a link must exist in order to explain the observed data when accounting for the error model of the noise. The ability to prove the existence of links under mild assumptions is in our opinion so valuable in knowledge generation that network models generated by other methods only should be used when the methods outperform RNI.

#### 4.2 When and why does LASSO fail?

The indicators SIC and WIC (Strong and Weak Irrepresentable Conditions) are fulfilled, *i.e.*  $\mu$  is below one, only for the data sets with a low condition number and SNR of one or higher, see Figure 5. This suggests that the response matrix, which is the transpose of the regressor matrix, needs to have a low condition number for accurate GRN estimation using LASSO. In our simulations LASSO typically fails due to introduction of several false positive links, Figure S.2.

Another indicator is  $r_{min}$ , the minimum ratio between the shortest regressor corresponding to a nonzero element and the longest corresponding to a zero. All data sets with a low condition number have a considerably higher  $r_{min}$  than all data sets with a high condition number, see Figure 6. The fact that  $r_{min}$  is below one for all data sets implies that the longest regressor corresponding to a non-existing link exceeds the length of the shortest regressor corresponding to an existing link. For the data sets with



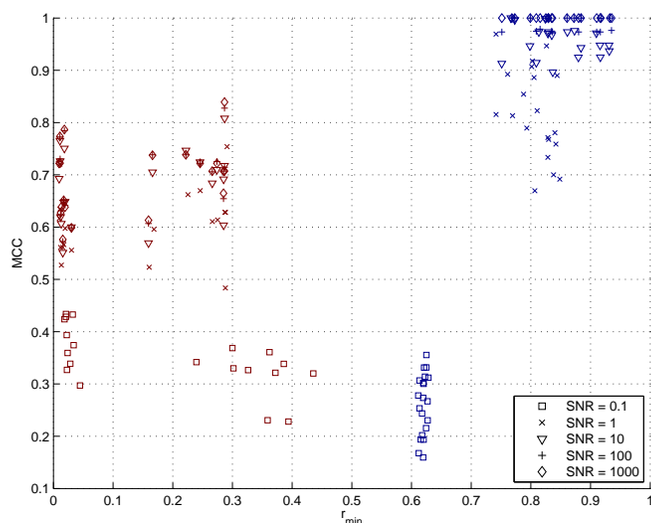
**Fig. 5** LASSO GRN inference accuracy versus irrepresentable condition. SIC and WIC are fulfilled only for the data sets with a low condition number (red) and an SNR of one or higher. The data sets with a high condition number (blue) all have a  $\mu$  above one.  $\mu$  describes the irrepresentable condition (10).

low condition number, the longest regressor corresponding to a non-existing link is expected to be nearly orthogonal to all regressors corresponding to existing links, while in data sets with high condition number they are not.

Evaluation of the irrepresentable conditions and ratio between the shortest regressor corresponding to a nonzero element and the longest corresponding to a zero require knowledge of the true network, so they cannot in practice be used to evaluate if LASSO will produce an accurate estimate. The lack of a linear relation between MCC and  $\mu$  or  $r_{min}$  indicates that neither of the measures captures all aspects that affect the performance of LASSO, so further studies are needed. Until a better testable criterion for failure of LASSO is presented, we recommend all users to check the condition number of the response matrix as discussed above. The condition number has the advantage of being a classical tool in linear algebra that is easy to calculate.

#### 4.3 Analysis of biological data

How well do the tested methods perform on real biological data? Although we can not control or vary the conditions of real data, we can take a data set and examine how well the methods can use it to infer a reference GRN. Such data has *e.g.* been collected by Lorenz *et al.*<sup>2</sup> for the Snf1 signalling pathway in *S. cerevisiae*, and they



**Fig. 6** LASSO GRN inference accuracy versus minimum ratio between the shortest regressor corresponding to a nonzero element and the longest corresponding to a zero over all rows. A clear separation between data sets where LASSO can infer the true structure (red) and where it cannot (blue) is seen at 0.5 for the ratio  $r_{min}$ . Colours as in Figure 5.

provide two reference GRNs. They perturbed the ten genes of the Snf1 signalling pathway in *S. cerevisiae* by inserting a plasmid containing an extra copy of each gene one-by-one and recording the resulting expression change of all ten genes. We calculated the weighted mean variance based on the reported propagated standard error of each data point of the response and perturbation matrix to 0.8 and 0.4, respectively. Because the variance of the response is twice as large as the variance of the perturbations, we calculate the  $SNR \approx 0.01$  using (8) with  $\alpha = 0.05$ . The closest data point based on the number of genes  $N = 10$ , estimated degree of interampattness 253<sup>32</sup>, the condition number  $\kappa(\mathbf{Y}) = 215$ , and SNR 0.01 in our simulated data is SNR 0.1 in Figure 1. Considering that (i) the estimated degree of interampattness is roughly twice that of the most interampatte simulated 10 gene network (108), (ii) the condition number is larger than the largest condition number of the simulated data sets (181), (iii) the number of perturbations are half, and (iv) the SNR is one order of magnitude lower, the expected MCCs should be well below 0.5. It is worth noting that the expected MCC of RNI is zero, *i.e.* the data is expected to contain so little information that no link can be proven to exist.

To avoid the influence of the selection of the regularisation parameter  $\zeta$  we varied it over the whole range from a full to an empty inferred network, see supplemental, and

**Table 5** Optimal performance based on MCC similarity with reported networks in Lorenz *et al.*<sup>2</sup>.

|             | S10  | S19  | S9   |
|-------------|------|------|------|
| LASSO       | 0.18 | 0.22 | 0.36 |
| LSCO        | 0.21 | 0.20 | 0.32 |
| Elastic net | 0.18 | 0.27 | 0.40 |
| RNI         | 0.00 | 0.00 | 0.00 |
| NIR (S9)    | 0.25 | 0.28 | 1.00 |

report the largest MCC value of each method for each of the networks reported by Lorenz *et al.*<sup>2</sup>–S10, S19, and S9. The first golden standard S10 is a collection of links that Lorenz *et al.*<sup>2</sup> found experimental evidence for in the literature. To get the second golden standard S19 they complemented these links with links that they found in their validation experiments using ChIP and qPCR. S9 is their final network estimate using NIR<sup>39</sup> followed by t-tests keeping only statistically significant interactions. It is not a gold standard but we included it for comparison. Note that a link in these golden standards can mean very different things; anything from a binding observed in a ChIP experiment to an influence on the expression of the other gene. The applied inference methods can only pick up influences that led to expression changes present in the recorded data. It is therefore unlikely that any of these golden standards equal the “true” network that would be achieved if more data was collected until Nordling’s confidence score for each possible link either is above one or approaches zero. We therefore refrain from making statements about which method that performs best based on comparison to these golden standards. The MCC of LASSO, Elastic Net, LSCO, and RNI is below 0.27 for both golden standards and hence in agreement with our expectation based on our simulations, see Table 5. The MCC between S9 and the two golden standards is below 0.28, *i.e.* of the same magnitude. The RNI inferred network is empty, indicating that the data contains so little information that no link can be proven to exist.

## 5 Conclusions

We have shown that all the tested  $L_1$  regularisation methods—LASSO, Elastic Net, and Bolasso—typically perform poorly in GRN inference when using data as ill-conditioned as typical experimental data. Testing on the *in vivo* data collected by Lorenz *et al.*<sup>2</sup> concurs with the expected performance in our simulations based on the properties of the data. As we use the regularisation coefficient that gives the most accurate network for each data set and noise realisation, the here reported perfor-

mance of LASSO, Elastic Net, Bolasso, and LSCO is in general unrealistically good. We can therefore with certainty say that the  $L_1$  regularisation methods fails for ill-conditioned data matrices even when the data is informative enough for network inference, while LSCO does not. However, this does not necessarily imply that LSCO in practice is always better and preferable over LASSO, Elastic Net, or Bolasso, because the accuracy is sensitive to the selection of the value of the regularisation coefficient<sup>26</sup>. Nonetheless, LSCO and RNI can yield better network estimates when the data is ill-conditioned so it is worth applying them. When the data is informative enough for network inference then all existing links can be proven to exist and RNI recovers the correct network structure<sup>32</sup>. As can be expected, we observed that LASSO fails when the SIC and WIC criteria are not fulfilled.

For both well-conditioned and ill-conditioned data, we found an SNR, as defined in (8), of 10 to be sufficient for LSCO and RNI to achieve maximum accuracies close to one. For data with an SNR below one the accuracy of all methods was in general low. This puts high demands on the quality of experimental data to be useful for GRN inference.

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