Network-Based Identification of Adaptive Pathways in Evolved Ethanol-Tolerant Bacterial Populations

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Abstract

Efficient production of ethanol for use as a renewable fuel requires organisms with a high level of ethanol tolerance. However, this trait is complex and increased tolerance therefore requires mutations in multiple genes and pathways. Here, we use experimental evolution for a system-level analysis of adaptation of Escherichia coli to high ethanol stress. As adaptation to extreme stress often results in complex mutational data sets consisting of both causal and noncausal passenger mutations, identifying the true adaptive mutations in these settings is not trivial. Therefore, we developed a novel method named IAMBEE (Identification of Adaptive Mutations in Bacterial Evolution Experiments). IAMBEE exploits the temporal profile of the acquisition of mutations during evolution in combination with the functional implications of each mutation at the protein level. These data are mapped to a genome-wide interaction network to search for adaptive mutations at the level of pathways. The 16 evolved populations in our data set together harbored 2,286 mutated genes with 4,470 unique mutations. Analysis by IAMBEE significantly reduced this number and resulted in identification of 90 mutated genes and 345 unique mutations that are most likely to be adaptive. Moreover, IAMBEE not only enabled the identification of previously known pathways involved in ethanol tolerance, but also identified novel systems such as the AcrAB-TolC efflux pump and fatty acids biosynthesis and even allowed to gain insight into the temporal profile of adaptation to ethanol stress. Furthermore, this method offers a solid framework for identifying the molecular underpinnings of other complex traits as well.

Key words: experimental evolution, biological networks, ethanol tolerance, bacteria, hypermutation, gene prioritization.

Introduction

Experimental evolution offers great potential to gain insights into the molecular mechanisms that contribute to the acquisition of complex traits (Kawecki et al. 2012; Wiser et al. 2013; Anderson et al. 2014). Previously, experimental evolution has been used not only to study the mechanisms underlying clinically (Palmer and Kishony 2013; Van den Bergh et al. 2016; Steenackers et al. 2016) or industrially relevant phenotypes (Winkler and Kao 2014), but also to improve key industrial traits for the production of advanced evolutionary engineered strains (Winkler and Kao 2014). Laboratory evolution experiments usually start from a single clone that is cultivated for prolonged periods of time in predefined conditions. During this period of time, natural selection favors mutations that confer a benefit in the chosen condition leading to improved phenotypes (Dragosits and Mattanovich 2013). Fitness is tracked over time and clones displaying increased fitness are genotyped to identify the underlying mutations (Schlöterer et al. 2015). Although some phenotypes are established by only one or just a few mutations, complex traits often involve mutations in multiple adaptive pathways, severely complicating identification of the causal adaptive mutations (Eyre-Walker 2010).

In this study, we used experimental evolution to study high ethanol tolerance in the bacterium Escherichia coli. Usually, microbial ethanol production capacity is severely limited by the toxic effect of ethanol itself. Therefore, higher ethanol tolerance and increased ethanol production are inherently linked (Huffer et al. 2012; Thammasittirong et al. 2013). Even though understanding and improving this trait is vital for strain engineering, it has been challenging to fully elucidate the underlying mechanisms. Previous studies have identified single genes (Gonzalez et al. 2003; Goodarzi et al. 2010) as well as epistatically interacting genes (Nicolaou et al. 2012)
involved in higher ethanol tolerance. However, tolerance to ethanol is clearly a complex trait established by the interaction of multiple genes and pathways (Gonzalez et al. 2003; Goodarzi et al. 2010; Nicolau et al. 2012; Swinnen et al. 2012; Lam et al. 2014) and a broad understanding of ethanol tolerance in E. coli is currently lacking. Moreover, in a previous study we found that hypermutation drives evolution under severe stress, such as ethanol stress, to enable adaptation of at least some individuals to avoid extinction (Swings et al. 2017). An increased mutation rate was found in all high ethanol tolerant populations and resulted in a higher ratio of passenger versus adaptive mutations, leading to an extremely complex mutational profile. This complexity is reflected in our data set of 16 evolved populations by a total of 2,286 mutated genes, containing 4,470 unique mutations. Consequently, the increased mutation rate combined with the complexity of the phenotype impedes the ability to statistically distinguish between true adaptive mutations and passenger mutations.

In most studies, this distinction between adaptive and passenger mutations is based on identifying mutations or mutated genes that recurrently emerge in independent evolutionary lines (Woods et al. 2006; Dees et al. 2012; Lawrence et al. 2013; Tamborero et al. 2013; Read and Massey 2014; Chen and Shapiro 2015; Hammerstrom et al. 2015). This narrow definition of parallelism assumes that only frequently mutated genes contribute to an adaptive phenotype. However, in populations that evolve independently, there is no guarantee that exactly the same mutation or even the same mutated gene is responsible for the observed adaptive phenotype. Affecting the same pathway through different and not necessarily frequently mutated genes might equally well induce the same adaptive phenotype (Tenaillon et al. 2012; Kvitk et al. 2013; Hong and Gresham 2014). Rather than identifying recurrent mutations, one can search for consistently mutated molecular pathways, assuming that adaptive mutations will hit the same adaptive pathways in independently evolved populations. Approaches that search for consistent changes in molecular pathways are typically network-based and have been applied successfully, mainly in the context of cancer genomics (Leiserson et al. 2015; Babur et al. 2015; Le Van et al. 2016; Pulido-Tamayo et al. 2016) but not yet for the mapping of genotypes to complex traits in clonal micro-organisms such as bacteria.

To cope with the specificities of clonal evolution experiments that aim to study complex traits, we developed a novel network-based method, IAMBEE. The method exploits the information gained from the trajectory of individual mutations along the evolution experiment together with network information to reduce the complexity of identifying adaptive pathways genes. Our experimental set-up combined with this unique network-based approach resulted in the identification of several adaptive pathways that conferred ethanol resistance in E. coli. The role of the 30S ribosomal subunit pathway (Mars et al. 2015; Suzuki et al. 2015) as well as the osmotic stress response pathway (ompR/envZ) (Cai and Inouye 2002; Quinn et al. 2014) were confirmed. In addition, newly predicted molecular mechanisms such as the multidrug efflux pump AcrAB-tolC and the fatty acid biosynthesis pathway were experimentally validated. These results demonstrate the value of IAMBEE to analyze complex mutational data sets, including even data sets resulting from a hypermutator phenotype, to obtain a comprehensive overview of the pathways and its specific mutated components involved in the establishment of a trait.

Results

Ethanol Tolerant Populations Display a Hypermutator Phenotype

We set up an evolution experiment in which 16 independent E. coli populations were experimentally evolved under increasing ethanol concentrations (fig. 1). Changes in ethanol tolerance due to accumulation of beneficial mutations were tracked over time to obtain a fitness trajectory for each population (supplementary fig. 1, Supplementary Material online). These trajectories for all 16 populations show remarkable selective sweeps between 5% and 6% ethanol tolerance (further referred to as the initial selective sweep) and from 6% to 6.5% ethanol tolerance (further referred to as the second selective sweep). The populations were sampled right before and right after each increase in ethanol tolerance and were subjected to pooled sequencing. Primary analysis of the data showed that each of the ethanol tolerant populations evolved a hypermutation phenotype. In depth study of this observation led to the conclusion that near-lethal conditions require rapid adaptation of at least some individuals to avoid extinction of the population (Swings et al. 2017). Hypermutation considerably facilitates rapid adaptation by increasing the probability to acquire a beneficial mutation (Chao and Cox, 1983; Taddei et al. 1997; Woods et al. 2011; Swings et al. 2017). Although hypermutation enables adaptation, it also leads to complex mutational profiles with multiple mutations in random genes (Woods et al. 2006), further impeding identification of causal mutations. In our data set of 16 evolved populations a total of 2,286 mutated genes, containing 4,470 unique mutations were detected. To identify causal mutation despite this mutational complexity, we developed a new method that overcomes the limitations of only identifying recurrent mutations in evolved populations.

Exploiting Parallel Evolution to Identify Adaptive Pathways

To distinguish between adaptive and passenger mutations and to identify pathways underlying complex traits we have developed IAMBEE, which integrates prior information on gene interactions (i.e., an interaction network) with the specificities of the experimental design. Key to the concept of IAMBEE is the use of multiple independently evolved populations to search for recurrently mutated molecular pathways. This search is driven by the interaction network and based on a decision theoretic subnetwork inference problem (De Maeyer et al. 2015; De Maeyer et al. 2016). However, given the high mutation rate and the relatively low number of independently evolved populations, additional information in the form of functional impact scores of the individual mutations is needed to drive
the analysis. We hereby assume a priori that not all mutations are equally likely to be involved in the adaptive phenotype. Mutations that increase in frequency during a selective sweep and/or that have a functional impact on the protein in which they occur, are more likely to be involved in the phenotype.

Figure 1 gives a conceptual overview of IAMBEE. The input consists of called mutations from multiple, independently evolved populations and a genome-wide interaction network of the organism of interest. After topology-weighting the interaction network to downweight the effect of hubs on the final solution (see Materials and Methods), IAMBEE proceeds in three steps (fig. 1): 1) The relevance score is calculated for each mutated gene in each population. The relevance score consists of three components. The first component describes the change in frequency of the mutation during a selective sweep in the population. Mutations that increase in frequency during a sweep are more likely to be adaptive than mutations that decrease in frequency. However, not necessarily all adaptive mutations will increase in frequency during a sweep (e.g., potentiating mutations) and conversely...
passenger mutations that hitchhike with driver mutations will also increase in frequency (Lang et al. 2013). Therefore, the frequency change component is complemented with a second component: the functional impact score. The functional impact score reflects the effect of the mutation on the function of the protein. Mutations that are likely to alter a protein’s function are more likely to be adaptive. A last component of a mutation’s relevance score relates to the mutation rate of the population in which the mutation occurs: we assume that mutations that originate from populations with a significantly higher mutation rate than the other populations should contribute relatively less to the final solution as they contain a larger number of passenger mutations (more noise) and a mutation of such a line should thus exhibit a stronger signal in order to be selected. A detailed overview of the calculation of the relevance scores by IAMBEE is shown in figure 2 and is described in the Materials and Methods. 2) The pathfinding step embodies the search for paths, which are defined as consecutive sets of edges connecting mutated genes from different populations, on the topology-weighted interaction network. These paths are weighted based on the relevance scores of the involved mutations and the weights of the edges involved in the path. The weight of a path reflects the degree of belief that the path is involved in the adaptive phenotype. 3) Subnetwork inference (optimization strategy) is subsequently used to select a subset of the paths found during the pathfinding step. This subset is selected in such a way that a maximum number of mutated genes with high relevance scores are included but a minimal number of edges is selected. This means that overlapping paths are more easily selected as they share edges, which reflects the search for molecular pathways that are consistently mutated throughout independently evolved populations. The resulting subset of paths makes up a subnetwork that consists of multiple connected components which are parts of molecular pathways. For a more detailed explanation of these steps, we refer to the Materials and Methods section.

**Fig. 2.** Calculation of the relevance scores for each mutation by IAMBEE. In the data acquisition step, a selective sweep of interest is chosen from an evolution experiment involving multiple parallel evolved populations. Samples taken at time points just before (blue arrows) and just after (orange arrows) this selective sweep are sequenced and mutations are called. For every mutation, a functional impact score and frequency change in the population are determined by the IAMBEE software. The frequency change is derived from the degree to which the mutation changes in frequency before (blue) and after (orange) the selective sweep. Genes with mutations that rise in frequency have higher frequency increase scores (green square) while a low frequency increase score is assigned to genes with mutations that decrease in frequency in the population (red squares). Next, a functional impact score is assigned to each mutation by using SIFT4G (Vaser et al. 2016). Genes with mutations having a high functional impact score are depicted with green triangles and vice versa. In addition, populations with a mutation rate that is significantly higher than the mutation rates of the other populations are detected. The relevance of mutated genes in populations with a significantly higher mutation rate are corrected (red star) to avoid overrepresentation of mutations from these populations. Finally, combining a gene’s frequency score, functional impact score and the correction for mutation rates allows calculating a relevance score for every mutated gene in every population. Mutated genes with a high relevance score (green circles) are more likely to harbor mutations that increase in frequency during the selective sweep, have high functional impact scores and are not involved in a population with significantly higher mutation rate than the rest of the populations.
Validation of IAMBEE Using Synthetic Data
To validate and characterize IAMBEE, we generated 100 synthetic data sets, each with randomly selected adaptive and passenger mutations (supplementary Material, Supplementary Material online). Running IAMBEE on these data sets resulted in one subnetwork containing prioritized mutated genes per combination of data set and parameter setting. Every synthetic data set was run with 50 different parameter settings ranging from settings that result in small subnetworks to settings resulting in larger subnetworks. As in this synthetic setting the true adaptive mutations (true positives) are known, we used PPV (the ratio of true positives to the total number of mutations prioritized by IAMBEE) and sensitivity (the number of true positives to the total number of true positives in the data set) as performance criteria. Results showed that, as expected for a method that makes relevant nonrandom predictions, small subnetworks have a high PPV at the expense of a lower sensitivity and subnetworks (of any size) rarely have both low sensitivity and PPV (supplementary fig. 2, Supplementary Material online). Exploring small solutions allows identifying a restricted number of candidates suitable for experimental validation, whereas exploring larger solutions provides a more complete pathway level view of the adaptive phenotype, but risks identifying false positives. The reliability of the predictions as estimated from the parameter settings at which they were detected is also included in the output of IAMBEE where more opaque edges represent edges which are involved in both small and large solutions (high PPV) whereas less opaque edges are only involved in small solutions (lower PPV) (figs. 3 and 4).

Network-Based Analysis Unravels Adaptive Pathways for High Ethanol Tolerance
We pooled the mutation data observed in the 16 different lines for, respectively, the first and second selective sweep and applied IAMBEE to the pooled data of each sweep to unveil pathways that drive increases in ethanol tolerance during each of the sweeps. We identified connected components that were common to both sweeps and components that were unique to each of the sweeps. Identified connected network components representative of adaptive pathways or at least parts of adaptive pathways are shown in figures 3 and 4, respectively, the initial and second selective sweep. During the initial selective sweep 32 connected network components, involving 108 genes harboring 228 mutations, were prioritized by IAMBEE out of a total of 1,666 mutated genes harboring 2,511 mutations in 16 populations. Likewise, in the second selective sweep 22 connected components, involving 90 genes harboring 345 mutations, were prioritized by IAMBEE out of a total 2,286 mutated genes harboring 4,470 mutations in the same 16 populations. 15 of these connected components were partly or entirely selected in both sweeps (supplementary fig. 3, Supplementary Material online). The fact that components were detected that are common to both the initial and second sweep but also specific for each of the sweeps demonstrates fundamental differences between initial adaptation to high ethanol stress and adaptation to prolonged exposure to increasing ethanol concentrations.

Below, we describe important identified connected network components and their putative roles in ethanol tolerance. A more detailed overview and description of all identified ethanol tolerance related pathways is given in the supplementary Results, Supplementary Material online.

The Fatty Acid Biosynthesis Pathway Is Selected Exclusively for Initial Adaptation
An important network component that was exclusively identified for the initial selective sweep corresponds to the fatty acid biosynthesis pathway, encoded by the fab genes. The prioritization of this pathway in the initial, but not in the second selective sweep, means that most mutations in the fab genes were already fixed in the latter step, explaining why they were not selected as being adaptive in the second selective sweep. This early fixation of fab mutations indicates that changing the fatty acid composition in the membrane is an initial adaptation strategy, but does not suffice to confer resistance to higher concentrations of ethanol. Different parallel populations accumulated mutations in both fabA and fabB. The amount of unsaturated fatty acids eventually present in the membrane depends on the competition for intermediates at the FabA branch point in the pathway (Heath and Rock 1996; Loffeld and Keweloh 1996; Zhang and Rock 2008). Mutations in either the fabA gene itself or in genes downstream (e.g., fabG or fabG) can permanently change the ratio of saturated versus unsaturated fatty acids thereby changing the fluidity of the membrane. Changes in membrane composition, such as the ratio of saturated versus unsaturated fatty acids, have previously been reported to affect ethanol tolerance (Burtke and Ingram 1980; Luo et al. 2009) although mutations in the fab genes have not been associated with ethanol resistance in the past.

To validate whether mutations in the fab genes affect membrane composition, we compared fatty acid content of selected strains harboring these mutations with that of the wild type (fig. 5). When exposed to 5% ethanol, the percentage of unsaturated fatty acids in wild-type cells increases 2-fold (fig. 5a). This observation corroborates previous results showing ethanol-induced inhibition of saturated fatty acid synthesis (Burtke and Ingram 1980). The percentage unsaturated fatty acids in the absence of ethanol in the two mutant populations harboring a fabA (HT15) and a fabB (HT12) mutation equals that of the wild-type ancestor. However, in the mutant populations subjected to 5% ethanol the percentage unsaturated fatty acids also increases, but not to the same extent as for the wild type (fig. 5b). This difference suggests a direct effect of the identified fab mutations on the ethanol-induced shift in saturated versus unsaturated fatty acids ratio. Increased proportions of unsaturated fatty acids fluidize the membrane. Mutations in the fab genes possibly counteract this shift to become more tolerant against ethanol by maintaining structural rigidity of the membrane.

Pathways Involved in Both Initial and Consecutive Adaptation
Several network components were prioritized by IAMBEE in, respectively, the first and second sweep that corresponded to
the same pathway and hence most likely to confer ethanol tolerance. When a putative adaptive pathway is selected in both sweeps, typically few mutated genes were detected in the initial sweep, whereas the remainder of the mutated genes was identified in the second sweep (figs. 3 and 4).

**Multidrug Efflux Pumps.** One network component of particular interest is linked to multidrug efflux complexes. The genes *acrA*, *acrB*, and *acrD*, encoding the AcrAB-TolC and the AcrAD-TolC multidrug efflux pump were found to be frequently mutated during the initial selective sweep.

Multidrug efflux pumps usually consist of three parts: an inner-membrane transporter, such as AcrB, a membrane fusion protein such as AcrA and an outer-membrane transport channel such as TolC. AcrD, like AcrB, binds to AcrA and forms a complex with TolC to constitute a multidrug efflux pump. Despite their association with tolerance to organic solvents (White et al. 1997; Atsumi et al. 2010; Bernardi et al. 2016), efflux pumps have to our knowledge not yet been specifically linked to ethanol tolerance. To validate the role of the efflux pump in ethanol tolerance, we constructed a
deletion mutation in $acrB$, one of the subunits of the AcrAB-TolC multidrug efflux pump that accumulated most mutations. Indeed, the $\Delta acrB$ deletion mutant has an increased growth rate compared with the wild-type ancestor under 5.5% ethanol stress (fig. 6). Moreover, the relative difference in growth compared with the wild type increases with the ethanol percentage in the medium (supplementary fig. 4, Supplementary Material online). Even though drug resistance mediated by efflux pumps is usually rendered by active efflux of the compound, our results suggest that ethanol tolerance through mutations in $acrAB$ works the other way around. Ethanol might be leaking into the cell through the efflux pump, consequently causing higher ethanol tolerance in an AcrAB deletion mutant. Previously, AcrAB-mediated organic solvent tolerance has been reported to depend on the polarity of the solvent, where deleting $acrA$ causes increased tolerance to the slightly polar isobutanol, suggesting a similar tolerance mechanism for ethanol which is even more polar (White et al. 1997; Atsumi et al. 2010).

Whereas $acrA$, $acrB$, and $acrD$ are mutated in the initial selective sweep, during the second selective sweep additional mutations occurred in $mdtA$ and $mdtF$. MdtA, like AcrA, is a membrane fusion protein in the MdtABC-TolC multidrug efflux pump (Borges-Walmsley et al. 2002; Nagakubo et al. 2002). MdtF is the counterpart of AcrB and acts as a transporter in the MdtEF-TolC multidrug efflux pump (Nishino and Yamaguchi 2002). The mutations in $mdtA$ and $mdtF$ that rise in frequency during a second selective sweep frequently occurred in populations that already harbored a mutation in the AcrAB-TolC efflux pump originating from the earlier selective sweep (four out of seven). Subsequent mutations in paralogous AcrAB-TolC and MdtABC-TolC multidrug efflux systems thus are likely to enable gradual adaption and further improve fitness under high ethanol concentrations. In addition, even though TolC is the common outer membrane channel for all above mentioned efflux pumps no mutations occurred in the $tolC$ gene. However, TolC is essential for cell functioning, suggesting that mutations in $tolC$ most likely have a high fitness cost compared with the advantage they can provide under ethanol stress.

Other network components were prioritized in both sweeps and have previously been associated with ethanol
This confirms the ability of IAMBBEE to identify true adaptive pathways. Below, we list the most relevant of these pathways. For a more in-depth description we refer to the supplementary Results, Supplementary Material online.

**DNA Repair.** One highly prioritized network component includes several genes involved in DNA repair mechanisms, such as the methyl-directed mismatch repair pathway (MMR, mutS, mutL, and mutH), the nucleotide excision repair pathway (NER, uvrA, uvrB, and uvrC) and the DNA helicase encoded by uvrD which is involved in both the MMR and NER pathways. Mutations in the aforementioned genes explain the observed higher mutation rates that were observed in the evolution experiment. Prioritization of this network component substantiates our previous work where we demonstrated the crucial role of mutations in the mismatch repair pathway for adaptation to ethanol under high stress conditions (Swings et al. 2017).

**Transcription and Translation.** Other highly prioritized network components are linked to transcription and translation. We found a frequently mutated connected network component containing several well-known genes involved in transcription termination and antitermination (rho, nusG, and nusA). Additionally, we found a frequently mutated subnet-work containing translation-linked genes encoding for 30 S ribosomal subunit proteins (rpsL, rpsD, rpsB, and rpsH) and the protein chain initiation factor infB. The Selection of genes involved in these pathways further corroborates earlier results that demonstrate early ethanol-induced transcription termination and ethanol-induced translational misreading during protein synthesis. Mutations in genes involved in transcription or translation might confer higher ethanol tolerance through compensation for these toxic effects (Burns and Richardson 1995; Pasman and von Hippel 2000; Haft et al. 2014).

Remarkably, mutations prioritized by IAMBBEE in, respectively, (anti) termination factors, such as rho, nusA, and nusG and ribosomal genes, such as rpsB, rpsD, rpsL, and rpsH tend to cooccur. These are consistently observed in the same populations and follow the same trajectories, suggesting that they are dependent on each other in driving the observed tolerance towards ethanol (figs. 3 and 4). This observation suggests
an epistatic interaction that corresponds to previous findings of Freddolino et al. who showed that mutations in *rpsl* increase ethanol tolerance in a mutant *rho* background (Freddolino et al. 2012) and Haft et al. (2014) who demonstrated the role of epistatic interactions between a mutation in *rho* and *rpsQ* in ethanol tolerance.

**Osmotic Stress Response.** The *envZ-ompR* two-component system that regulates outer membrane porin genes in response to changes in extracellular osmotic pressure was also prioritized (Cai and Inouye 2002; Quinn et al. 2014). Ethanol increases membrane fluidity, thereby causing leakage and osmotic stress as shown by Goodarzi et al. (2010). Mutations in *envZ* and *ompR* suggest adaptations to the increased osmotic stress under high ethanol conditions.

**Amino Acid Biosynthesis.** Finally, several of the prioritized network components and genes are involved in amino acid biosynthesis, such as isoleucine and valine biosynthesis (*ilvD, ilvC, ilvB, ilvE, ilvL*, and *tdcB*), alanine and phenylalanine biosynthesis (*ilvE* and *alaA*), methionine biosynthesis (*metE, metB, and metH*), biosynthesis of tetrahydrofolic acid, which is a precursor in the metabolism of amino acids (*purH*), a gene involved in arginine biosynthesis (*argH*), and a gene involved in threonine and glycine biosynthesis (*itaE*) (Keseler et al. 2013). A role of amino acid biosynthesis and transport in tolerance towards ethanol stress has previously been suggested in yeast, because of impaired delivery of amino acids into the cell as a result of membrane functions being disrupted by ethanol (Yoshikawa et al. 2009; Stanley et al. 2010).

**Pathways Exclusively Involved in the Second Adaptation Step**
One smaller network component that was exclusively but highly prioritized in the second selective sweep consists of two genes *fadB* and *gabT*. FadB plays a role in fatty acid oxidation and is regulated by FadR (DiRusso et al. 1992). Interestingly, a *fadR* deletion mutant was also recently found to increase organic solvent tolerance (Oh et al. 2012) pointing to a similar process.

Several additional identified network components correspond to pathways that have never been linked to ethanol tolerance before, but that might influence this trait are discussed in the supplementary Results, Supplementary Material online. In conclusion, we state that IAMBEE is able to detect previously known as well as new adaptive pathways. The identified adaptive pathways in ethanol tolerance might serve as a basis for future strain improvement efforts.

**Indications for Epistasis at the Pathway Level**
Remarkably, adaptive mutations in the fatty acid pathways tend to occur in a mutual exclusive way: eight populations have mutations in *fadB-gabT*, nine populations have mutations in the FabA-B system, two populations (HT13 and HT14) have mutations in both pathways and only one population (HT3) has no mutations in either of the respective pathways. Strikingly, this would imply negative epistasis at the pathway level, i.e., a mutation in either pathway increases ethanol resistance but mutations in both mechanisms do not lead to a greater increase in resistance. The incomplete pattern of mutual exclusivity in HT14 can be explained by the fact that both the mutation in *fabA* (present in 50% of population) and *fadB* (present in 12% of population) are not fixed in the population. Therefore, it is possible that these mutations exist in different subpopulations. As was the case in the cooccurrence of mutations in the AcrAB-toIC/fab pathway, the incomplete pattern of mutual exclusivity in HT13 could be explained by the fact that the *fabA* mutation is situated towards the end of the fabA protein (seven amino acids near the end), which makes it likely that this mutation is not functionally relevant. Indeed, by determining the membrane composition we confirmed that the percentage of unsaturated fatty acids in absence and in response to 5% ethanol did not differ from the wild-type strain, suggesting that this particular mutation does not contribute to higher ethanol stress by changing the membrane composition (supplementary fig. 5, Supplementary Material online).

Additionally, mutations in the AcrAB-ToIC efflux pump tend to cooccur with mutations in the previously mentioned *fab* pathway. From the ten populations which have a mutation in *acrab-tolC* and the nine populations which have a mutation in the *fab* pathway, eight populations overlap. According to the mutational trajectories, mutations in both pathways arise during the same selective sweep (5–6%) in seven populations. Only in one population (HT10), the mutation in *acrab-tolC* was obtained late during the second selective sweep, following an earlier mutation in the *fab* pathway. Only population HT13 had a *fabA* mutation but not a mutation in *acrab-tolC*. Again, this specific mutation in *fabA* is located at seven amino acid residues near the end of the FabA protein (supplementary fig. 5, Supplementary Material online).

**Comparison with a per Gene Mutation Frequency Approach**
To show the value of the network-based approach of IAMBEE, we compared our results with those obtained by a frequency-based approach which ranks genes based on the number of populations in which they were mutated (fig. 7, supplementary file 1, Supplementary Material online). Although IAMBEE does not explicitly search for genes that are recurrently mutated across populations, results show that IAMBEE is able to also prioritize most of the frequently mutated genes which are associated with ethanol resistance (e.g., *rpsl* and *envZ*). Exclusively mapping frequently mutated genes (136 and 153 for, respectively, the first and the second selective sweep) to the genome-wide interaction network showed (supplementary figs. 6 and 7, Supplementary Material online) that adaptive pathways which were identified by IAMBEE, such as the fatty acid biosynthesis pathway, fabA-gabT and AcrAB-ToIC, are largely or completely missed (supplementary figs. 6 and 7, Supplementary Material online). This can be explained by the fact that these pathways are composed of genes that are not necessarily frequently mutated (e.g., some connecting genes were only found mutated in one or two populations). By exploiting all mutated genes over the network and using information from mutational...
trajectories and functional impact scores IAMBE can, in contrast to the frequency-based approach, extract adaptive pathways consisting of less frequently mutated but highly connected genes (supplementary table 1, Supplementary Material online). However, as IAMBE is network-based it is possible that some genes are missed because they are not present in the network (e.g., marC and tsaA [Atsumi et al. 2010; Minty et al. 2011]). Nevertheless, this does not outweigh the benefit of IAMBE to retrieve more complete pathways and to enable reasoning about the temporal aspects of mutation acquisition.

**Discussion**

Evolution experiments have been successfully used to identify the role of specific genes in an adaptive phenotype (Suzuki et al. 2014; Van den Bergh et al. 2016). However, genetic data derived from parallel evolution experiments is usually interpreted by looking at the mutation frequencies of the individual genes (Woods et al. 2006). Especially when dealing with complex traits and hypermutation, these studies do not necessarily yield insight into the complex interactions of the genes that contribute to the adaptive phenotype. Key to unraveling the genetic mechanisms underlying high ethanol tolerance is the development of a dedicated analysis method. IAMBE is unique in prioritizing adaptive mutations by combining information on each individual mutation inferred from functional impact scores and relative frequency increases during a selective sweep, with information on the interactions between genes (a genome-wide interaction network). Using our newly developed method, we were able to prioritize multiple pathways that were recurrently mutated in different independent high ethanol tolerant populations. Among the highly prioritized pathways, those related to translation, anti-termination and amino acid metabolism were previously associated with high ethanol tolerance. Recovering these well-known pathways confirms the ability of IAMBE to identify true adaptive pathways.

On top of those well-known systems, we identified a yet undescribed role for multidrug efflux pumps in the continuous adaptation to high ethanol stress and the role of fatty acid metabolism in allowing the cell to cope with the toxic effects of ethanol on the membrane (Dombek and Ingram 1984). Related to the latter mechanism, binding and penetration of ethanol into the lipid bilayer increases membrane fluidity (Ramos et al. 2002), thereby inducing secondary effects, such as osmotic stress (Ingram and Buttke 1984; Ingram 1990; Huffer et al. 2011). Response to osmotic stress has previously been shown to induce alterations of the membrane composition including cis-to-trans isomerization of unsaturated fatty-acids as a short-term response (Loffeld and Kewloho 1996) and alteration of the ratio of saturated versus unsaturated fatty acids as a long-term response (Buttke and Ingram 1980; Luo et al. 2009). Both changes to the membrane composition can result in denser packing of the fatty acids thereby increasing the rigidity of the membrane which enables the cell to withstand the toxic effect of ethanol (Ramos et al. 2002; Nicolau et al. 2012). Although there has been confusion about the effect of unsaturated fatty acids on ethanol tolerance (Buttke and Ingram 1980; Luo et al. 2009; Vanegas et al. 2012), we provide evidence that tempering the shift to higher ratios of unsaturated fatty acids confers higher resistance to ethanol (fig. 5). We could indeed show that mutations in representative genes of the fab pathway resulted in increased ethanol tolerance by affecting the ratio of saturated versus unsaturated fatty acids.

As IAMBE is designed to be used in combination with a dedicated experimental set-up, which includes sequencing of the evolving populations before and after a selective sweep, it is possible to gain insight in the temporal profile of adaptation. Using this unique feature of the method we found that mutations in the fatty acid biosynthesis pathway occur early in the evolutionary trajectory of ethanol resistance in E. coli whereas mutations in other pathways have less strict temporal constraints. In this context, we also found that mutations in the pathway for fatty acid biosynthesis (fabA, fabG, and fabB) and a pathway involved in fatty acid oxidation (gabT, fadB) were mutually exclusive while having high coverage (15 out of the 16 populations had a mutation in either pathway). This implies that fatty acids play a pivotal role in ethanol tolerance, but that either a mutation in one of the two pathways does not lead to a significant increase in fitness if a mutation in the other pathway is already present (negative epistasis) or that having a mutation simultaneously in both pathways is lethal (synthetic lethality as an extreme form of negative epistasis). In contrast, mutations in the fatty acid
biosynthesis pathway (fabA, fabG, and fabB) and the AcrAB-TolC efflux pump (acrA and acrB) significantly co-occur, suggesting positive epistasis between these pathways.

When compared with a naive approach which is based on recurrence of mutations across experiments at the level of individual genes, it is obvious that IAMBEE offers not only the advantage of being able to identify adaptive mutations which are not frequently mutated but also to interpret adaptive mutations and epistasis at the level of pathways. As such, IAMBEE is very useful and meets the need for adequate tools to analyze highly complex mutational data sets.

Conclusions

Experimental evolution can readily yield insight in complex traits assuming low complexity of the resulting mutational profiles. However, in the case of complex traits and especially when hypermutation arises, evolution experiments often lead to complex mutational profiles with high rates of passenger mutations that are difficult to interpret. Traditionally, adaptive genes are identified by counting the number of mutations per gene across independently evolved populations. Although this approach is valid in some data sets, in complex mutational profiles it neglects less frequently mutated genes, resulting in the inability to generate a broad understanding of the adaptive phenotype. Therefore, we developed IAMBEE, a method that exploits the interaction network, combined with information from mutational trajectories and functional impact scores, to identify adaptive genes and pathways from complex mutational data sets. By applying IAMBEE to an evolution experiment consisting of 16 independently evolved E. coli populations subjected to increasing ethanol concentrations, pathways that were previously linked to high ethanol resistance as well as novel pathways, which were experimentally validated, could be identified. In conclusion, IAMBEE is a powerful tool that successfully allows to generate a broader understanding of (complex) traits that could not be fully elucidated so far.

Materials and Methods

Data Acquisition

Experimental Evolution

We used a set of 16 parallel evolved E. coli populations. Of these highly ethanol-tolerant populations, seven originated from our previously conducted evolution experiment (Swings et al. 2017) and we initiated a new experiment using the same workflow to add an additional nine populations to the final data set. All populations acquired a hypermutator phenotype, which is necessary to enable evolution under near-lethal stress conditions (Swings et al. 2017). For ease of reading, we renamed all populations HT1-16 (High Tolerance). In brief, all parallel populations originated from the same ancestral strains SX4 and SX25. We used lysogeny broth (LB) supplemented with 5% (v/v) ethanol as primary stress conditions (Swings et al. 2017). For ease of phenotype, which is necessary to enable evolution under the final data set. All populations acquired a hypermutator (Swings et al. 2017) and we initiated a new experiment using from our previously conducted evolution experiment

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Mutation Calling

High-quality genomic DNA from overnight cultures of the ancestor and intermediate points of evolved populations was isolated (DNeasy Blood and Tissue kit, Qiagen). 100 bp paired-end sequencing libraries with an average insert size of 200 bp were prepared at GeneCore (EMBL, Heidelberg) and used for massive parallel sequencing with the Illumina HiSeq2000. We used CLC Genomics Workbench version 7.6 (https://www.qiagenbioinformatics.com) (RRID: SCR_011853) for analysis of the sequences. Following quality assessment of the raw data, reads were trimmed using quality scores of the individual bases (quality limit = 0.01; maximum number of ambiguous bases = 2). Reads shorter than 15 bases were discarded from the set. We used the CLC Assembly Cell 4.0 algorithm to map the trimmed reads to the E. coli MG1655 reference genome (NC_000913.1) yielding a minimal coverage of 150× (mismatch cost = 2; insertion cost = 3; deletion cost = 3; length fraction = 0.8; similarity fraction = 0.8). Mutations were called using the CLC Low Frequency Variant Detector (http://resources.qiagenbioinformatics.com/manuals/clccancerresearchworkbench/200/index.php? manual=Low_Frequency_Variant_Detection.html; last accessed August 3, 2017; required significance = 1%; minimum coverage = 10; minimum frequency = 10%). Finally, the mutations in the SX4 compared with the MG1655 reference genome were discarded. In addition, CLC largely fails in the detection of larger rearrangements, such as large insertions, deletions, inversions or duplications. Therefore, we performed an additional analysis on the sequence data using a dedicated tool, Pindel (Ye et al. 2009), for detection of larger genomic rearrangements. We ran Pindel using default settings.

Mapping of Mutations to Genes

Mutations in coding regions of genes were mapped to those genes, whereas mutations in intergenic regions were only mapped to a gene if they resided in a promoter. Promoter element locations were taken from RegulonDB (RRID: SCR_003499) (Gama-Castro et al. 2016) and PromBase
(Kanhere and Bansal 2005). Supplementary table 3, Supplementary Material online, shows the number of mutations retained and used in both analyses.

**Functional Impact Scores and Frequency of Mutations**

To calculate functional impact scores for each mutation we used SIFT scores which were calculated using the SIFT4G annotator version 2.2 with the *E. coli* (GCA_000005845.1.21) database (Vaser et al. 2016). Note that while SIFT scores were used in this paper, any functional impact score measure can be inserted in IAMBEE.

As IAMBEE tries to identify the causal molecular pathways which lie at the basis of a selective sweep, mutations which decrease in frequency during a selective sweep are not taken into account. Therefore, we have to determine when a mutation “decreases” in frequency. As the precision of frequency calling of mutations is finite, the naive way of viewing all mutations with a negative frequency increase as “decreasing” is not valid. This would discard too many mutations which remained stable or, most of the time, were fixed previously in the population. As these mutations could be potentiating mutations, we do not want to discard them. Because of this, and the specifications of the CLC variant caller (a required significance of 1%), we viewed all mutations with a decrease in frequency of at least 2% as decreasing. All mutations in both selective sweeps, together with their SIFT scores and their increase in frequency during the selective sweep are given in supplementary file 1, Supplementary Material online.

**Genome-Wide Interaction Network**

We used a directed genome-wide interaction network of *E. coli* K-12 MG1655 compiled from (de)methylation, (de)phosphorylation and metabolic interactions from KEGG version 80 (RRID: SCR_012773) (Kanehisa et al. 2014; Kanehisa et al. 2016), protein–DNA, sigma factor binding and sRNA–DNA interactions from reguLonDB version 9.2 (Gama-Castro et al. 2016) and protein–protein interactions from STRING version 10 (RRID: SCR_005223) (Szklarczyk et al. 2015). To reduce the number of false positive interactions in the interaction network, only direct (physical) associations with a score of at least 0.8 were retained from STRING. Interactions involving the primary sigma factor RpoD were removed as RpoD regulates over half of the genes in the interaction network. Furthermore, self-edges were deleted. The final genome-wide interaction network contains 2,678 nodes (genes) and 14,702 edges (interactions between genes/sRNAs), representing about 63% of *E. coli* K-12 genes. This interaction network is supplied together with IAMBEE at http://bioinformatics.intec.ugent.be/IAMBEE.

**Construction of the Probabilistic Genome-Wide Interaction Network**

IAMBEE is guided by a directed genome-wide interaction network with the nodes representing genes and the edges representing interactions between these genes. A topology-based weighting of the genome-wide interaction network was performed to reduce the effect of hubs in the subsequent analysis steps: a power law distribution (Barabási and Albert 1999) was estimated based on the out-degrees of the nodes in the interaction network. Next, a sigmoidal function was constructed using as inflection point the out-degree that corresponded to the 90th percentile. This leads to following topology-based weighting of each edge between node i and node j (De Maeyer et al. 2015):

\[
\text{weight}(ij) = \frac{1}{1 + e^{\frac{\text{out}\_\text{degree}(i) - \text{out}\_\text{degree}(j)}{\text{inflection}\_\text{point}}}}
\]

This sigmoidal function is utilized to mainly down weight interactions originating from large hubs while avoiding to penalize interactions involving nodes with low out-degrees.

**IAMBEE**

**Calculation of Relevance Scores**

Not all mutations are equally likely to be involved in the adaptive phenotype. Therefore, a relevance score was assigned to each mutation based on its estimated functional impact on the coding/promoter sequence and based on its relative increase in frequency in the population during a fitness increase. The functional impact score reflects how likely a mutation causes a functional change in the resulting protein(s). Here it is based on the degree of conservation of amino acid residues in sequence alignments from closely related sequences using the SIFT algorithm (RRID: SCR_012813) (Ng and Henikoff 2001; Kumar et al. 2009; Vaser et al. 2016). To derive frequency increases, for each population the adaptive trajectory (e.g., fitness profile) is used to delineate selective sweeps (sudden jumps in fitness or an increase in adaptation towards the experimental conditions). The frequency increase of a mutation is equal to the difference of its frequency in the population just after and just before the sweep. To assess the relative importance of a mutation’s frequency increase or functional impact score, we first estimate both the distribution functions, based on the frequency increase/impact score of all mutations from all evolved populations. As neither the functional impact score distribution, nor the frequency increase distribution is expected to follow any known mathematical distribution, the distributions are estimated using a nonparametric cumulative distribution function (MathWorks, Inc. 2017). As synonymous mutations would skew the distribution of the functional impact scores towards low functional impact scores, which could result in assigning relatively high relevance scores to mutations with poor functional impact scores, synonymous mutations are removed from the data when estimating the functional impact distribution function. Note that because some synonymous mutations do have relevant functional impact scores they are not discarded but only ignored when estimating the functional impact distribution. Although the functional impact distribution function is estimated using all mutation data from all evolved populations, the frequency increase distribution is estimated on a per-population basis as the population dynamics can differ between populations. This means that one functional impact distribution is estimated, whereas the number of frequency increase distributions is
equal to the number of parallel evolved populations are estimated.

Based on these distributions a relevance score is calculated for each mutated gene in each population as follows:

$$
\text{relevance}(S, n) = (1 - eCDF_{fun}(\text{Functional}_\text{score}(S, n))) \\
\times (eCDF_{freq}(\text{Frequency}_\text{increase}(S, n)))
$$

with $eCDF_{fun}(\text{Functional}_\text{core}(S, n))$ the value of the cumulative distribution function of the functional impact scores for the mutation in gene $S$ in population $n$ with the most deleterious functional impact score (note the $1-eCDF_{fun}$ as we used SIFT scores and a low SIFT score corresponds to a high functional impact), $eCDF_{freq}(\text{Frequency}_\text{increase}(S, n))$ the value of the cumulative distribution function in population $n$ of the frequency increases for the mutation in gene $S$ in that population with the highest frequency increase. $\text{relevance}(S, n)$ is a value between 0 (gene $S$ is unlikely to be relevant towards adaptation in population $n$) and 1 (gene $S$ is very likely to be relevant towards adaptation in population $n$). Genes without mutations are assigned the mean functional impact score and frequency increase when calculating their relevance.

Furthermore, if the data set contains populations with a mutation rate which is significantly higher than the mutation rates of the other populations, the search for paths in the pathfinding step (see following paragraph) would be skewed towards this population (supplementary table 1, Supplementary Material online). To reduce the impact of the population’s mutation rate on the pathfinding without completely discarding these populations, a correction factor for each population is calculated. To detect populations with significantly higher rates we use the modified $Z$-score for outlier detection (Iglewicz and Hoaglin 1993) as follows:

$$
\text{modified } Z \text{ score}(n) = \frac{0.6745 \times (\text{mutations}(n) - \text{median}(n_1, \ldots, n_i))}{\text{MAD}(n)}
$$

with $\text{MAD}(n) = \text{median}(|\text{mutations}(n) - \text{median}(n_1, \ldots, n_i)|)$

with $\text{mutations}(n)$ the number of mutations in population $n$, $\text{median}(n_1, \ldots, n_i)$ the median number of mutations in a population, and $\text{MAD}(n)$ the mean absolute deviation of population $n$. Note that in the original publication the modified $Z$ score is defined as the absolute value of the measure used in this paper. We intentionally left out the absolute value to avoid down weighting populations with few mutations. Populations with a significantly higher mutation rate are defined as populations having a modified $Z$ score of at least 3.5 (Iglewicz and Hoaglin 1993). From this modified $Z$ score a population specific correction factor is calculated, based on a parameter $p$ which sets the upper limit for the correction factor. In our analysis, we set this to 3 to have an upper limit of 0.85 but based on how a user would like to deal with populations having significantly higher mutation rates, the factor can be anywhere between 0 and 3.5:

$$
correction(n) = \begin{cases} 
\frac{P}{\text{modified } Z \text{ score}(n)} & \text{if modified } Z \text{ score}(n) \geq 3.5 \\
1 & \text{else} 
\end{cases}
$$

Due to the modified $Z$ score, the correction factor intrinsically assigns a lower value to outlier populations if a larger number of independent populations are available, hereby largely reducing the effects of populations with high mutation rates to reduce noise when a large number of independent populations is present. When only a limited number of independent populations is available, the correction factor will be higher as in that case populations with larger mutation rates are needed to exploit parallelism.

The relevance score and the correction factor are integrated into a single score for every mutated gene in every population. This is implemented as follows:

$$
\text{corrected relevance}(S, n) = \text{relevance}(S, n) \times \text{correction}(n)
$$

with $S$ a mutated gene in population $n$.

**Pathfinding between Mutated Genes**

All genes with at least one mutation in any independent population are mapped on the topology-weighted genome-wide interaction network. Subsequently, all possible paths originating from a mutated gene in a population and ending in any other gene which is mutated in another population, are enumerated. A path is defined as a series of consecutive edges in the interaction network. We exclude paths between mutated genes in the same populations, reasoning that because of the clonality a single mutation in a pathway will confer most of its fitness advantage (Pulido-Tamayo et al. 2016) and including paths between mutated genes within one population would not be informative as this does not reflect parallel evolution.

Each path is assigned a probability which reflects the degree of belief that the path is associated with the adaptive phenotype under study. This probability takes into account the weights of the edges which make up the path (calculated based on the network topology in the previous step) and the corrected relevance scores from both the start gene and the terminal gene of the path (calculated based on the frequency increase and the functional impact score of both genes in the data preparation step). Only the relevance scores of the start gene and the terminal gene are considered irrespective of whether or not intermediate genes are mutated. If intermediate genes would be taken into account, even one passenger mutation in the middle of an interesting molecular pathway would severely decrease the probability of every found path in that molecular pathway. This leads to the following equation for the probability of a path:
probability(S, n, E, m)_{S\neq E, n\neq m} = \prod_{(i, j) \in P} (weight_{(i, j)})^{*relevance(S, n)*relevance(E, m)}

with (S, n, E, m) the path which starts in gene S, which is mutated in population n and terminates in gene E, which is mutated in population m. P is the collection of edges which make up the path and (i, j) is the edge from node i to node j.

Enumerating all possible paths is computationally expensive and leads to a prohibitively large computational cost in the subsequent subnetwork inference step. Therefore, the following heuristics are used: 1) Based on biological considerations (Gitter et al. 2011; Navlakha et al. 2012) the maximum path length is set to four. 2) From all possible paths originating from a mutated gene in a specific population, only the 25 paths with highest probabilities are retained.

Subnetwork Inference and Prioritization of Molecular Pathways

The final step of the analysis is the inference of a subnetwork containing the molecular pathways responsible for the adaptive phenotype. This subnetwork consists of a subset of the paths selected in the previous step. This subset of paths is obtained by optimizing the following function:

\[ S(K) = \sum_{n \in R} \left( \sum_{S \in Q_n} (P(path(mut_{S,n}, mut_{all})|probabilities)) \right) - |K|x_e \]

Where S(K) is the score of the selected subnetwork and needs to be maximized, |K| is the number of edges selected, x_e is the imposed cost for each edge, R is the collection of populations used in the experiment, Q_n is the collection of mutated genes from population n and P(path(mut_{S,n}, mut_{all})|probabilities) is the probability that there exists a path relevant to the observed phenotype between a mutated gene S in population n and any other mutated gene in any other population, given the degrees of belief (probabilities) of all found paths in the path-finding step. The calculation of this term is a generalization of the two-terminal reliability problem (Fratta and Montanari 1973; Cook and Ramirez-Marquez 2007). Note that the optimal subnetwork, which is the selected subnetwork with the highest score S(K), is not necessarily a connected graph.

As the complexity of this problem inhibits a deterministic solution, a greedy hill-climbing heuristic is used in which the previously found paths get sampled pseudo-randomly based on the overlap the paths have with each other. Overlapping paths are more likely to be sampled together. As this procedure is stochastic in nature, the procedure is repeated 20 times and the best solution with respect to the optimization score S(K) is used as the solution.

The x_e parameter is an important parameter as it incentivizes IAMBEE to primarily select overlapping paths with high probabilities because doing so a single edge can be used multiple times while the cost for selecting this edge only has to be paid once. This is biologically relevant as molecular mechanisms in which multiple (partly) overlapping paths with high probabilities are found, are likely mechanisms of interest for a specific selective sweep.

Setting the x_e parameter is not trivial as its optimal value is data set specific. If x_e is set too high the subnetwork will be small and multiple causal molecular pathways are likely missed. Conversely if x_e is set too low the subnetwork will be too big and of little practical use as the fraction of false positives in the solution increases (supplementary fig. 2, Supplementary Material online). Therefore, instead of calculating the optimal subnetwork for one specific cost, we perform a parameter sweep over the x_e parameter and summarize the results in the form of a network, which is obtained by taking the union of all found optimal subnetworks and where the edges are prioritized based on the maximum edge cost for which they are still included in an optimal subnetwork. This means that edges with a high priority (visualized in the output as opaque edges) get selected even when the edge cost x_e is high. This is useful as the PPV (positive predictive value) of a subset of opaque edges is higher (Synthetic data in supplementary Material, Supplementary Material online) and thus a good starting point for experimental validation.

Parameter Setting

The parameters of IAMBEE were set as follows for both jumps in ethanol tolerance: The path length was kept at the default value of 4 and the maximum number of paths between every pair of mutated genes was kept at the default value of 25. The sweep over the edge cost parameter x_e was set from 0.1 to 1.5 in steps of 0.025 and the maximum size for an optimal subnetwork to be accepted was set to 80 (in terms of nodes) in order to keep the resulting subnetwork small enough to interpret manually.

Validation of IAMBEE Features

To show the importance of the different features of IAMBEE, which include mutation frequencies, functional scores, a correction factor for populations with extreme mutation rates and the use of an interaction network, the method was adjusted several times to exclude one feature each time. Each of these adjusted versions of IAMBEE was applied to the synthetic data set. PPV and sensitivity plots were constructed (supplementary fig. 2, Supplementary Material online) and demonstrated that each feature led to an increase in performance. The results are discussed in supplementary Material, Supplementary Material online.

Biological Validation Methods

The methods for the biological validation experiments can be found in the supplementary Methods section, Supplementary Material online.

Availability of Data and Material

The genome sequencing data set generated and analyzed during the current study are available in the SRA repository of NCBI, PRJNA380734 (https://www.ncbi.nlm.nih.gov/bioproject/380734; last accessed June 15, 2017).
IAMBEE is available at http://bioinformatics.intagec.ugent.be/IAMBEE.

Supplementary Material
Supplementary data are available at Molecular Biology and Evolution online.

Author Contributions
T.S. and B.W. conceptualized the study, analyzed and interpreted the results and wrote the manuscript. T.S. designed and performed the biological experiments. B.W. designed IAMBEE and used it to analyze the sequence data. T.S. and C.B. helped in performing the biological experiments. N.V., J.M., and K.M. conceptualized the study, designed the experiments, discussed the results, and edited the manuscript.

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