Impact of Chromosomal Inversions on the Yeast DAL Cluster

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Abstract
Chromosomal rearrangements occur readily in nature and are a major reshaping force during genome evolution. Such large scale modifications are usually deleterious causing several fitness defects, but sometimes can confer an advantage and become adaptive. For example the DAL metabolic cluster in yeast was assembled in recent evolutionary times in the Hemiascomycetes lineage, through a set of rearrangements that brought together the genes involved in the allantoin degradation pathway. In eukaryotes, the existence of physical clustering of genes with related functions supports the notion that neighbouring ORFs tend to be co-expressed and that the order of genes along the chromosomes may have biological significance, rather than being random as previously believed. In this study, we investigate the phenotypic effect that inversions have on the DAL gene cluster, expressed during nitrogen starvation. In all Saccharomyces “sensu stricto” species the order of the DAL cluster is conserved, while in the “sensu lato” species Naumovia castellii, which grows significantly worse than S. cerevisiae on allantoin, the cluster includes two nested inversions encompassing three DAL genes. We constructed several inverted and non-inverted S. cerevisiae strains possessing different inversions including those to mimic the configuration of the N. castellii DAL cluster. We showed that the inversion of DAL2 lower its own expression and reduces yeast fitness during nitrogen starvation. This rearrangement also altered the expression of the neighbouring genes DAL1 and DAL4. Moreover, we showed that the expression of the DAL4 anti-sense transcript (SUT614) does not change upon inversions of DAL2 and therefore is unlikely to be involved in its regulation. These results show that the order of the DAL cluster has an impact on the phenotype and gene expression, suggesting that these rearrangements may have been adaptive in the “sensu stricto” group in relation to the low availability of nitrogen in the environment.

Introduction
The first evidence of chromosomal inversions was published in 1921 by Sturtevant who studied rearrangement of genes in Drosophila [1]. The current literature shows that inversions have been found in almost all the organisms ranging from prokaryotes to eukaryotes and its rate vary among different lineages [2,3,4,5,6]. Chromosomal inversions in yeast are predominantly small sometimes including only one gene and can generate a new gene order putting the relevant loci in proximity. Such small chromosomal inversions may be partially responsible for the separation of Candida albicans and Saccharomyces cerevisiae, since 1,100 single-gene inversions have occurred since the divergence of the two species [7].

It is important to study inversions as they can affect the gene expression of the inverted genes as well as of the neighbouring genes. Genome studies provide an increasing evidence that expression and regulation of genes is not only controlled independently by its own promoters and associated regulatory elements but is also dependent on its location in the genome [8,9,10]. It has been indicated that similarly expressed genes are clustered in genomic neighbourhood in almost all taxa, however, it is so far not clear what mechanism enforces them to be clustered or co-localized in the genome. In prokaryotes cluster of genes form operons leading to a strong co-expression and in eukaryotes physically linked genes are co-expressed. The formation of co-expressed clusters during the period of evolution further promotes the fact that gene expression is dependent on its position in the genome [11,12,13,14]. In eukaryotes, it has been argued that co-regulation may not be needed for the physical clustering of genes, and that evolutionary pressure could be sufficient to bring functional genes together [15,16].

In yeast, Saccharomyces cerevisiae, the genes involved in the mitotic cell cycle were the first to be shown as co-expressed clusters in the genome [17]. The size of co-expressed clusters in yeast is relatively small, not exceeding ten genes or few kilobases compared to the clusters of multicellular eukaryotes consisting of 20–30 genes [18].

There are two large groups of gene clusters in S. cerevisiae, the GAL cluster [19] and the DAL cluster [20]. The DAL cluster is the largest metabolic gene cluster, enabling yeast to use allantoin as a non-preferred nitrogen source. Allantoin is converted through a series of steps into ammonia which is a simpler form of nitrogen readily used by yeast (Figure 1A). The genes in this cluster were initially scattered throughout the yeast genome and co-localised only recently within a single sub-telomeric site on chromosome IX in the ancestor of S. cerevisiae and N. castellii. The DAL cluster in S. cerevisiae consists of six adjacent genes encoding for six of the eight proteins involved in allantoin degradation. While the gene order of
the DAL cluster is completely conserved in the *Saccharomyces* “sensu stricto” species, in *N. castellii*, belonging to the “sensu lato” group, the cluster includes two nested inversions (Figure 1B). Other more distantly related hemiascomycetes species still maintain the DAL genes on different chromosomes. DAL5 (located on chromosome X and encodes for allantoate permease) and DUR1,2 (located on chromosome II and encodes for a protein which is responsible for converting urea to ammonia) are the two genes which are not located in the DAL cluster [21].

Computational studies provide us with the evidence that species belonging to the *Saccharomyces* “sensu stricto” group present different orientation of the DAL genes compared to *N. castellii* [21]. However, the potential role of these inversions on the phenotype has not been tested experimentally. In this study, we set to investigate the effects of different inversions of the DAL cluster on the gene expression and yeast growth. We constructed various inverted and non-inverted control strains possessing single, double or triple gene inversion as well as a strain resembling the *N. castellii* DAL structure in *S. cerevisiae* background.

We analyzed the fitness of the wild type, inverted and non-inverted control strains in allantoin containing medium that is responsible for triggering the expression of the DAL genes. We showed that the strain mimicking *N. castellii* DAL cluster possessed a lower growth rate as compared to the non-inverted control and the wild type strains. Moreover, we showed that the DAL2 inversion alone was responsible for the fitness impairment. Gene expression data indicated that DAL4 gene was down regulated the most in the DAL2 inverted strain background while the DAL4 antisense level did not change. Overall, our results suggest that the rearrangements of the DAL cluster in the *S. cerevisiae* “sensu stricto” species may have conferred some advantage in natural environment knowingly low in nitrogen.

Figure 1. Schematic representation of the yeast DAL gene cluster involved in the allantoin degradation pathway. The panel A shows the six DAL genes located on chromosome IX at the same position and orientation in all *S. cerevisiae* “sensu stricto” species. In *N. castellii* the cluster presents two nested inversions involving DAL1, DAL2 and DAL4 (marked with a double ended arrow). The colour orange indicates collinear genes, the pale green arrows shows genes which underwent one inversion event and the dark green colour indicate the DAL2 gene which inverted twice in *N. castellii*. (Figure adapted from Wolf, 2006). The panel B shows how allantoin is converted to allantoate by allantoinase and degraded to produce glyoxalate and urea. In the final stage of the pathway, the glyoxalate is converted to malate and the urea to ammonia.

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Materials and Methods

Strains and Media
All strains were engineered with the FY3 background and they were maintained on YPD medium containing 2% (w/v) yeast extract, 1% (w/v) peptone and 2% (w/v) glucose. The transformants were grown on YPD-agar containing the desired antibiotics i.e. 300 μg/ml geneticin (GibcoBRL), 100 μg/ml cloNAT ( Werner BioAgents, Jena, Germany), 10 μg/ml phleomycin (Invivo-Gen) and 300 μg/ml hygromycin B (Duchefa Biochemic) for selection of the kanMX, natNT2, pCre-ble and hphNT1 markers. A full list of engineered strains is provided in Table S5. Mineral salt medium (F1 medium) was prepared as previously described [22]. 0.0125% (w/v) allantoin and 0.1% (w/v) proline+0.0125% (w/v) allantoin were used as nitrogen source.

Primers and Oligonucleotide probes
Gene sequences were obtained from SGD (http://www.yeastgenome.org/) and PCR primers were designed using the Primer3 programme. Strand specific oligonucleotide probes were manually designed keeping the GC content to 40% and length to 34–40 bp. The BLAST tool of SGD was used to check the specificity of each probe and primer. Primer and probe sequences are provided in Tables S1, S2, S3, S4.

Construction of Inverted and Non-inverted strains
The resistance gene marker cassettes used in this study were loxP-kanMX-loxP [23], loxP-hphNT1-loxP and lox2272-natNT2-lox2272 [24]. The loxP-kanMX-loxP cassette was amplified according to Delneri et al. 2003 [25] and loxP-hphNT1-loxP and lox2272-natNT2-lox2272 cassettes were amplified according to Janke et al. 2004 [26]. These cassettes were inserted in the genome of S. cerevisiae by PCR-mediated gene replacement method to construct our inverted and non-inverted strains [27]. Transformation was done by lithium acetate protocol [28]. The strains bearing cassettes were then transformed with Cre-recombinase containing plasmid. The Cre-recombinase enzyme was induced by first growing the cells overnight in YP-rafainose medium and then in YP-galactose for 2–3 hrs. The colonies were verified for inversion and non-inversion by colony PCR [25]. All the primers used for construction of strains are provided in Tables S1 and S4.

Fitness Growth Rate Assay
Growth rate of all the inverted and non-inverted strains was determined using FLUOstar optima microplate reader. Cells were grown to stationary phase in YPD, minimal and allantoin containing medium. Optical density (OD) of the cultures was measured at 395 nm and then diluted to OD395 nm = 0.1 in pre-warmed respective medium. 240 μl of the diluted cultures was transferred to each well of 96 well plate including the media controls. The OD measurement was taken by the microplate reader at 30°C for 40 hours at intervals of 5 minute and with 1 minute linear shaking just before every measurement. Growth curves were plotted using the Optima data analysis programme and standard deviations were calculated in Excel. The growth parameters (growth rate μ and maximum cell biomass A) were measured using R statistic package grofit [29].

RNA extraction and reverse transcription
Total RNA for reverse transcription was extracted using the Qiagen RNA extraction kit following the manufacturer’s instructions. RNA concentration was determined using the nanodrop spectrophotometer (ND-1000), quality and integrity of RNA was checked by electrophoresis on 1.5% (w/v) agarose gel. Whenever required RNA was treated with DNaseI (Fermentas) prior to cDNA synthesis as described by the manufacturer. 1 μg of total RNA was reverse transcribed to cDNA in a 20 μl reaction mixture by Qiagen reverse transcription kit using the random primers.

Gene expression analysis by real-time quantitative PCR
The expression levels of DAL1, DAL2 and DAL4 were determined using real-time PCR. All real time PCRs were performed on the cDNA samples using the Quantitect real time PCR kit from Qiagen on Chromo4 gradient thermocycler in 96 well plate from Biorad. Real time PCR primers were synthesized by MWG-Eurofins (HPSF purified). Each primer was designed to amplify a 250–300 bp fragment. Optimized reactions were carried out in 50 μl final volume containing 10 ng/μl of cDNA, 5 pmole of each primer and 25 μl of 2× quantitect syber green. The qPCR conditions were used with an initial denaturation of 3 min at 95°C followed by 35 cycles consisting of 95°C for 45 sec, 58°C for 45 sec and 72°C for 3 min with a final extension of 5 min at 72°C. Melting curves were analyzed from 55°C to 95°C at a rate of 0.2°C / 2 sec. Actin (ACT1) was used as a housekeeping reference gene. Serial dilutions (10−1–10−5) of actin DNA was used for generating standard curve. The expression of each gene was estimated using the Ct Values. All real time PCRs were tested in triplicate and each experiment was done on three independent biological replicates. The overall standard deviation is shown in results. A blank (with no RT) was also included in each experiment.

Northern hybridization
Total RNA for northern blotting was extracted from yeast strains using Trizol (Invitrogen, catalogue # 155-96-018) as described by the manufacturer. 20 μg of the total RNA was loaded in each slot and resolved on 1% (w/v) formaldehyde agarose gel [30]. Samples were prepared by adding RNA loading dye from Fermentas which contained ethiduim bromide. The RNA was transferred on the nylon membrane with panther semidry electroblotter HEP-1 in accordance to the manufacturer’s manual using 1XTBE as transfer buffer. The RNA was fixed on the membrane by UV irradiation for 1 min using the UV crosslinker (XL-1500) 1200 μl/cm2 for 60 sec. 5 pmole oligo was labelled with [32p] ATP using T4 polynucleotide kinase (Fermentas, catalogue #EK003). A mixture of three and five oligonucleotide probes binding at different regions in the gene were used for Actin and DAL4 respectively. The membrane was hybridized at 37°C in oligo hyb solution (0.17Moles Na2HPO4, 0.079Moles NaH2PO4, 35g SDS, 1 ml 0.5 M EDTA, dH2O to 500 ml and warm to dissolve, pH ~7.2) for overnight. The membranes were washed at 42°C in 6×SSC for 10 min and 2×SSC, 0.1% SDS for 10 min. The final washing was done at room temperature in 6×SSC. Membranes were exposed to phosphoimager screen for 1–3 days and band intensities were quantified using Quantitect programme from Biorad.

Results and Discussion

DAL cluster in S. cerevisiae and N. castelli
Naumovia castelli DAL cluster differs from Saccharomyces cerevisiae and other sensu stricto species by two nested gene inversion. We compared the sizes of the DAL cluster and that of the intergenic regions between S. cerevisiae and N. castelli (Figure 2). The overall size of the cluster was almost identical (only 1 base pair different) while the length of the intergenic regions seemed to vary between the two species. We also observed that the sizes of DAL1 and DAL3 differ significantly between S. cerevisiae and N. castelli. DAL1 is 7% longer in N. castelli than in S. cerevisiae and DAL3 is 29% smaller in

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N. castellii than in S. cerevisiae. Moreover, in S. cerevisiae there is an ARS sequence between DAL2 and DCG1. We did not find any ARS sequence in N. castellii DAL cluster.

Growth assays of S. cerevisiae and N. castellii in allantoin medium

The DAL cluster is expressed under nitrogen limited conditions to degrade allantoin (non-preferable source of nitrogen) to a simpler form of nitrogen such as ammonia. Previous literature have shown that DAL4 mutants grow normally in medium containing either ammonia, allantoate, arginine and asparagines as nitrogen sources, however, they are incapable to grow on allantoin containing medium [31].

Allantoin is a purine derivative formed by oxidation of xanthine to urate and then to allantoin in purine degradation pathway. This reaction is catalyzed by xanthine dehydrogenase (XDH) and urate oxidase (UOX). XDH is not present in yeast species and therefore they import urate, allantoin or allantoate into the cell to use purine derivatives as nitrogen source. The yeast species which do not possess DAL cluster are capable of using urate as a nitrogen source using urate permease (UAP) to import it and UOX to oxidize it and the enzymes of DAL metabolic pathway to degrade it to urea [32]. S. cerevisiae and N. castellii do not have UAP and UOX and therefore are unable to use urate and thus import allantoin as nitrogen source using allantoin permease gene DAL4. Once imported allantoin is then degraded to simpler form of nitrogen using the same DAL pathway genes in all yeasts however, in S. cerevisiae and N. castellii the DAL genes have been organized into a cluster. The ability to import allantoin instead of urate ruled out the oxygen requiring step performed by UOX leading to the biochemical reorganization of purine degradation pathway. The species which possess the DAL cluster have the ability to grow under oxygen limiting conditions [21].

Firstly, we tested the growth of our wild type strain on different concentrations of allantoin and found that a high concentration of 0.4% (w/v) was deleterious to the strain while a concentration of 0.0125% (w/v) was just suboptimal and therefore ideal to detect fitness changes in the engineered mutant strains (Figure S1). Then, we measured the growth rate of the wild type N. castellii and S. cerevisiae (FY3) in F1 medium and F1 medium containing allantoin as nitrogen source. It was observed that wild type N. castellii was growing much slower than S. cerevisiae in both allantoin and F1 media (Figure S2). There was a 7% decline of growth rate for N. castellii in F1 medium and 68% decline of growth rate in allantoin medium relative to S. cerevisiae. These results indicated that S. cerevisiae can utilize allantoin in much more growth efficient way than N. castellii wild type strain.

Construction of the S. cerevisiae strains with different gene inversions

To study the impact of gene inversions on gene expression, we constructed inverted and non-inverted control strains using the cre-loxP system [33]. Wild type S. cerevisiae (FY3) was the background strain used to construct all inverted and non-inverted control strains. The non-inverted control strains will be referred as “control” in the text. In total, we constructed two strains with a single gene inversion, one with double gene inversion, and two triple gene inversions along with their respective control strains which were collinear to S. cerevisiae but with two loxP or lox2272 scars at the inversion breakpoints (Figure S3). The lists of all the strains created in the work with the respective inversions are listed in Figure S3.

Compared to S. cerevisiae, the three genes DAL1, DAL2 and DAL4 are inverted in N. castellii, with DAL2 being inverted twice. Since it is not defined knowledge which gene got inverted first we adopted two different strategies to re-create the N. castellii like DAL cluster in S. cerevisiae background.

In the first strategy, the DAL2 gene was inverted first creating the DAL2.1 strain which was then used to generate the inversion of the DNA sequence containing DAL1, DAL4 and DAL2. In the
second strategy the DNA segment with DAL1, DAL4 and DAL2 was inverted first creating the DAL1-4-2.I2 strain, which was then used to invert the DAL2 gene. Both strategies resulted in the construction of a final S. cerevisiae strain with the DAL cluster order similar to that of N. castellii (DAL1-4-I2.I1 and DAL1-4-I2.I2) (Figure 3). Control strains were created identical to their parent strains except for the presence of the loxP and lox2272 scars (CI1, CI2, CII1 and CII2) (Figure 3).

Fitness assays of the engineered strains in different nitrogen sources

The fitness of inverted and control strains was determined in rich medium (YPD), minimal medium containing allantoin as nitrogen source and F1 medium. We did not observe any significant changes in the growth rate of all the inverted and control strains in YPD and F1 medium.

For the strains engineered using the first strategy, we observed that in the allantoin containing medium, the DAL2 gene inversion affected the growth rate of DAL2.I1 in comparison to the FY3 and CI1 strains. There was a 38% drop in growth rate for the DAL2.I1 strain relative to the CI1 and FY3 strains (Figure 4). The same loss of fitness was observed after the inversion of the DNA sequence containing DAL1, DAL4 and DAL2 in the DAL1-4-I2.I1 when compared to FY3 and CI1. This result suggested that the second larger inversion did not contribute any further to the defective phenotype.

Using the inverted strains created with the second strategy the strain DAL1-4-I2.I2 with the larger inversion encompassing DAL1, DAL2 and DAL4 did not show any change in fitness compared to the control and wild type strains in allantoin medium (Figure S4). However, when the second inversion of just DAL2 was generated in the DAL1-4-I2.I2 strain there was a drop in the growth rate of the inverted strains compared to the control and FY3 (Figure S4).

Again, these results suggest that the DAL2 inversion is the main cause of the drop in fitness. The other engineered inverted strains DAL3.I and DAL3-7.I did not show any significant gain or loss in the growth rate (Figure S5).

A recent study showed that clustering of GAL genes in S. cerevisiae is not linked to positive fitness effects, since the disruption of GAL gene cluster (GAL1-GAL10-GAL7) did not have any phenotypic effect [34]. Our study established that disrupting the gene order of the DAL metabolic cluster has an effect on the growth rate. In particular, the inversion of DAL2 was responsible of the fitness change.

Effect of DAL2 inversion on the expression of DAL cluster genes

We looked at the expression level of the DAL genes between inverted and control strains by performing real-time quantitative PCR. Any change in expression occurring due to insertion of loxPs or lox2272s was taken into consideration by normalizing the expression of inverted strains with the control strains. For each sample, the housekeeping ACT1 was amplified and all data were normalized to ACT1.

Our fitness assays showed that inversion of DAL2 reduces the growth rate of inverted strain as compared to the control strain. Our next approach was to study the effect of inversion on DAL2 expression and also of its neighbouring genes expression such as DAL1 and DAL4. We observed about 50% decline of DAL1 gene expression in the DAL2.I1 strain, while the DAL2 and DAL4 expression level was reduced to ca. 90% (Figure 5). The expression of DAL4 and DAL1 in control strains was similar to the WT, while the expression of DAL2 in the control strain was at least 50% lower than the wild type suggesting an effect of the loxP insertion in the transcript level. However, since no phenotypic effect was seen in Figure 3. Strategies followed to engineer strains possessing N. castellii like DAL cluster. In strategy one the DAL2 was inverted first followed by the larger inversion of the three genes DAL1, DAL4 and DAL2. In strategy two the block of the three genes, DAL1, DAL4 and DAL2, was inverted first followed by the single inversion of the DAL2 gene. The control strains had the loxP and lox2272 insertions in the intergenic regions, but they do not present the inversion. The red, yellow and green blocks indicate the single inverted, double inverted and non-inverted genes respectively. The loxP and lox2272 scars are represented as blue and violet triangles respectively.

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the control strain, the amount of \textit{DAL2} gene product must have been sufficient to grant normal growth.

Both DAL2.I1 and DAL1-4-I2.I2 has reduced expression of \textit{DAL4} in allantoin containing medium, and the reduced fitness effect could be explained by the down regulation of \textit{DAL4} as this gene is responsible for allantoin uptake by the cell.

Co-expression of genes can be due to the presence of bidirectional promoters [35], such as in the case of \textit{GAL1} and \textit{GAL10} genes in the GAL cluster [34], or by regulatory mechanisms at chromatin level [36]. Eukaryotic genes that are co-regulated in a cluster are usually inter dependent regarding their expression [37,38], and it has been reported that genomic neighbourhoods can regulate transcript levels, playing an important role in genome evolution [39,40]. Our results demonstrate that \textit{DAL2} inversion not only affects its own gene expression but also that one of neighbouring genes (Figure 5). Moreover, \textit{DAL4}, \textit{DAL1} and \textit{DCG1} (but not \textit{DAL2}) have an antisense transcript associated with their mRNA which may play a role in their regulation (Figure S6). Interestingly, studies on the impact of neighbourhood continuity on gene expression in engineered inverted and the non-inverted \textit{Drosophila} strains found that no significant changes occurred in the expression of neighbourhood genes, showing that cluster organisation may not be as important as previously thought in maintaining the correct level of mRNAs in the cell [41]. This difference outcome could be due to the fact that the intergenic regions in yeast are very short with a high density of tightly regulated functional units and therefore small changes in the sequence are more likely to cause an alteration of the gene expression compared to \textit{Drosophila}.

To investigate whether the expression of \textit{DAL4} antisense transcript, \textit{SUT614}, has changed upon inversion of \textit{DAL2}, we carried out strand specific Northern Blot analysis.

**Expression analysis of \textit{DAL4} and \textit{SUT614}**

Co-expression of closely located genes is not only dependent upon transcription factors or chromatin structure but is also

![Figure 4. Fitness assay of wild type (FY3) and engineered strains.](image_url)
dependent upon bidirectionally active promoters [42]. *Saccharomyces cerevisiae* possess several transcripts which either exist as stable unannotated transcripts which are SUT and CUT. We used the bidirectional promoter database developed by Xu et al, 2009 to find out the antisense transcripts for DAL genes [35]. It was observed that three out of six DAL genes in the cluster possess stable unannotated transcripts which are SUT195, SUT614 and SUT196. SUT614 is the antisense transcript of DAL4. To test the role of bidirectional promoter in DAL4 we looked at the expression of SUT614 and DAL4 in our DAL2.I1 and control strain. We therefore checked the expression of DAL4 and SUT614 in YPD medium and after inducing the DAL cluster in allantoin+proline containing medium. As expected, we observed different levels of expression for DAL4 and SUT614 in the two different media. As expected, we observed that in both CI1 and DAL2.I1 DAL4 is not expressed in YPD medium, while the antisense transcript showed a strong expression level (Figure 6A). The expression analysis of DAL4 and SUT614 in proline+allantoin medium showed that the DAL4 expression is reduced in DAL2.I1 when compared to its control CI1. Moreover, SUT614 was equally highly expressed in both strains (Figure 6B). This data shows that inverting DAL2 reduces the expression of its neighboring gene DAL4 but not that one of the antisense transcript, suggesting that SUT614 is not responsible for DAL4 regulation.

**Figure 5.** Expression levels of DAL2, DAL4 and DAL1 genes in the DAL2.I1 and CI1 strains. Real time PCR to show the expression of DAL1 (A), DAL2 (B) and DAL4 (C). Blue, green and red, boxes represents the FY3 control and inverted strains, respectively. Error bars are from three technical replicas for each of the three independent biological samples. Relative normalized fold expression was calculated by using ΔΔCt method and ACT1 as a reference gene. doi:10.1371/journal.pone.0042022.g005

**Figure 6.** Northern analysis of DAL4 sense and antisense transcripts. RNA was extracted from cells grown under non-induced conditions (YPD medium, panel A) and induced conditions (F1 medium containing Proline+Allantoin as N-source, panel B). Oligonucleotides specific for the DAL4 sense and anti-sense strand were used. As expected no expression of DAL4 sense transcript was observed in YPD medium while, the antisense DAL4 signal was very strong (A). Under induced conditions the expression of DAL4 sense was greatly reduced in the inverted strain as compared to the control while the expression of antisense transcript remained same in both strain backgrounds (B). *ACT1* was used as the reference gene for expression comparisons. doi:10.1371/journal.pone.0042022.g006
Conclusions and Future Prospects

In this work we re-created in S. cerevisiae the two nested inversions present in N. castellii DAL cluster, and we showed that the inversion of DAL2 gene alone was sufficient to impair the growth rate of this yeast when allantoin is given as source of nitrogen. Further studies on the effect of the DAL2 inversion in N. castellii background would be interesting to understand if the change in the gene order has an impact in the allantoin utilisation in this species as well.

In S. cerevisiae, the inverted DAL2.I1 strain affects its own gene expression and that of the neighbouring genes DAL1 and DAL4.

Since DAL2 inversion is changing the expression of DALA and DALI1, it is possible that DAL metabolic pathway is regulated by a feed-back mechanism involving the DAL2 product. Allantoin permease encoded by DALA is responsible for entry of allantoin into the cell, which is converted to allantoate with help of enzyme allantoinase encoded by DALI gene. DAL2 encodes for allantoicase and is the main central gene responsible for converting allantoate to simpler form of nitrogen through series of steps. Therefore, it is plausible that the impairment of DAL2 expression is affecting the whole DAL metabolic pathway and that with the down regulation of DALA the cell is unable to take-in sufficient allantoin for normal growth.

We showed that the inversion of DAL2 does not affect the expression of “SUT614”, although the DALA transcription is significantly compromised. It is therefore unlikely that DALA is regulated by its anti-sense transcript SUT614.

The DAL cluster is sensitive to nitrogen catabolite repression (NCR) and is expressed in the lack of readily available nitrogen sources. Its regulation occurs via three types of cis-acting and trans-acting factors: the UASNTR binding either Gln3p or Gat1p [43], present upstream of all allantoin pathway genes, the URSOGATAA associated with DAL80p [44] which is involved in the down-regulation of DAL gene expression, and the UISALL binding DAL81p and DAL82p [45]. As summarised in Figure S6, DAL1, DAL2 and DAL4 require functional GLN3, DAL82 and DAL81 for transcription. Although the molecular modifications carried out to construct the inverted and control strains did not disrupt specifically the transcription factors binding sites, it is possible that a change of DNA sequence in this location may hamper the ability of these transcription factors to dock on the DNA.

Moreover, the sub-telomeric DAL cluster is placed within Htz1-activated domain (HZAD) which consists of histone H2A variant H2A.Z (Htz1). H2A.Z is present in region of activated domain (HZAD) which consists of histone H2A variant H2A.Z (Htz1). H2A.Z is present in region of DAL metabolic cluster. It is thought to act as an anti-silencing factor by preventing the spread of Sir protein and these genes.

Supporting Information

Figure S1  Fitness assay to optimise allantoin concentration. Growth rate of a wild type S. cerevisiae strain, FY3, was measured in different concentrations of allantoin containing medium over the course of 40 hours. Higher concentration of allantoin 0.4% (w/v) (blue line), was found to be toxic to the cells. For the fitness assays a sub-optimal concentration of 0.0125% (w/v) of allantoin (red line) was used. Each point represents the mean average of 5 technical replicates. Error bars at 95% confidence interval. (TIF)

Figure S2  Fitness assay of N. castellii (WT) and FY3 in F1 and F1+allantoin medium. The growth profiles of S. cerevisiae and N. castellii was measured in F1 and F1+allantoin medium. N. castellii (red line) is less fit than S. cerevisiae (blue line) in both media (A). The maximum growth rate and cell biomass were calculated using the R statistic package grofit [46]. Each point represents the mean average of three technical replicas for five independent biological samples. Error bars are at 95% confidence intervals. (TIF)

Figure S3  The DAL cluster structure of wild type, inverted and control strains. The loxP sequences were inserted in FY3 strain at the inversion breakpoints to construct the single, double and triple inverted strains using the cre-lox system (A). The control strains without the inversions but carrying the loxP insertions are shown in panel B. The red and green blocks indicate the inverted and collinear genes, respectively, whereas the blue triangles represent the loxP scars. (TIF)

Figure S4  Fitness assay of wild type (FY3) and engineered strains. The growth rate of DAL1-I2-I2 strain (red line), FY3 (blue line) and control strain (green line) in F1 medium supplemented with 0.0125% (w/v) allantoin was found to be the same (A). The inversion of DAL2 in the DAL1-I2-I2 strain showed a drop in the growth rate of the inverted strains (yellow line) compared to the control strains (b). The growth curves were derived from the OD values obtained from plate reader. The error bars represent the mean of three technical replicates of five independent biological replicates for each strain respectively. Error bars are at 95% confidence intervals. (TIF)

Figure S5  Fitness assay of DAL3.I and DAL3-7.I along with their respective control strains. The growth rate of DAL3.I strain (red line), FY3 (blue line) and control strain (green line) in F1 medium supplemented with 0.0125% (w/v) allantoin was found to be the same (A). The DAL3-7.I inverted strain (red line) also possessed equal growth rate as compared to the FY3 (blue line) and control strain (green line). The growth curves were derived from the OD values obtained from plate reader. The error bars represent the mean of three technical replicates of five independent biological replicates for each strain respectively. Error bars are at 95% confidence intervals. (TIF)

Figure S6  The map of the transcription binding site and anti-sense transcript in the DAL cluster. Representation of the DAL genes sense transcripts (green arrows), anti-sense transcripts (blue arrows), sites of loxP insertions (red arrows) and transcription factor binding sites (black arrows). (TIF)

Table S1  List of checking primers used for confirming inverted and non-inverted strains. (DOC)

Table S2  List of primers used for real time PCR. (DOC)

Table S3  List of probes used for northern blotting. (DOC)

Table S4  The set of cassette amplifying primers for engineering inverted and non-inverted strains. (DOC)

Table S5  The genotypes of Saccharomyces cerevisiae inverted and non-inverted strains used in this study. (DOC)
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References

1. Sturtevant AH (1921) A case of rearrangement of genes in Drosophila. Proc Natl Acad Sci U S A 7: 235–237.
2. Bonierbale MW, Plaisted RL, Tankesley SD (1988) RFLP maps based on a common set of clones reveal modes of chromosomal evolution in Potato and Tomato. Genetics 120: 1095–1103.
3. Feder JL, Roethele JB, Flichak K, Niedbalski J, Romero-Severson J (2003) Evidence for inversion polymorphism related to sympatric host race formation in the apple maggot fly, Rhagoletis pomonella. Genetics 163: 939–953.
4. Shaw DD, Wilkinson P, Coates DJ (1983) Increased chromosomal mutation rate after hybridization between two subspecies of grasshoppers. Science 220: 1165–1167.
5. Stefansson H, Helgason A, Thorleifsson G, Steinthorsdottir V, Masson G, et al. (2005) A common inversion under selection in Europeans. Nat Genet 37: 129–137.
6. Warner JW (1976) Chromosomal variation in the plains woodrat: male meiosis and breeding studies. J Mammal 57: 10–18.
7. Sgroi G, Federspiel N, Jones T, Hansen N, Bivolarovic V, et al. (2000) Prevalence of small inversions in yeast gene order evolution. Proc Natl Acad Sci U S A 97: 14453–14457.
8. Germain HJ, Ingedam MH, Koster J, Goetz S, Seppen J, et al. (2007) Domain-wide regulation of gene expression in the human genome. Genome Res 17: 1296–1299.
9. Lee TI, Young RA (2000) Transcription of eukaryotic protein-coding genes. Annu Rev Genet 34: 77–132.
10. van Driel R, Franz F, Verschure PJ (2003) The eukaryotic genome: a system regulated at different hierarchical levels. J Cell Sci 116: 4067–4075.
11. Lercher MJ, Blumenthal T, Hurst LD (2002) Coexpression of neighboring genes in Caenorhabditis elegans is mostly due to opossums and duplicate genes. Genome Res 13: 238–243.
12. Lercher MJ, Urrutia AO, Hurst LD (2002) Clustering of housekeeping genes provides a unified model of gene order in the human genome. Nat Genet 31: 180–183.
13. Kruglyak S, Tang H (2000) Regulation of adjacent yeast genes. Trends Genet 16: 109–111.
14. Hurst LD, Pal C, Lercher MJ (2004) The evolutionary dynamics of eukaryotic gene order. Nat Rev Genet 5: 299–310.
15. Yerushalmi U, Teicher M (2007) Examining emergence of functional gene clustering in a simulated evolution. Bull Math Biol 69: 2261–2280.
16. Carter Z, Delneri D (2010) New generation of loxP-mutated deletion cassettes for the genetic manipulation of yeast natural isolates. Yeast 27: 765–773.
17. Dehler D, Hammarsten S, Holmgren S, Hopf I, Hultberg T, et al. (2003) Evidence for inversion polymorphism related to sympatric host race formation in the apple maggot fly, Rhagoletis pomonella. Genetics 163: 939–953.
18. Shaw DD, Wilkinson P, Coates DJ (1983) Increased chromosomal mutation rate after hybridization between two subspecies of grasshoppers. Science 220: 1165–1167.
19. Gietz RD, Schiestl RH (2007) High-efficiency yeast transformation using the LiAc/SS carrier DNA/PFG method. Nat Protoc 2: 31–34.
20. Kahn M, Hasenbrink G, Lichtenberg-Frate H, Ludwig J, Kuchichino M (2010) Genetic diversity and breeding studies. J Mammal 57: 1–18.
21. Pal C, Hurst LD (2003) Evidence for inversion polymorphism related to sympatric host race formation in the apple maggot fly, Rhagoletis pomonella. Genetics 163: 939–953.
22. Baganz F, Hayes A, Farquhar R, Butler PR, Gardner DC, et al. (1998) Quantitative analysis of yeast gene function using competition experiments in continuous culture. Yeast 14: 1417–1427.
23. Dekker U, Heath S, Felder T, Benhaim J, Hegenmann JH (1996) A new efficient gene disruption cassette for repeated use in budding yeast. Nucleic Acids Res 24: 2519–2524.
24. Carter Z, Delneri D (2010) New generation of loxP-mutated deletion cassettes for the genetic manipulation of yeast natural isolates. Yeast 27: 765–773.
25. Dehler D, Hammarsten S, Holmgren S, Hopf I, Hultberg T, et al. (2003) Evidence for inversion polymorphism related to sympatric host race formation in the apple maggot fly, Rhagoletis pomonella. Genetics 163: 939–953.
26. Shaw DD, Wilkinson P, Coates DJ (1983) Increased chromosomal mutation rate after hybridization between two subspecies of grasshoppers. Science 220: 1165–1167.
27. Wach A, Brachet A, Pohlmann R, Möller E (1994) New modular tools for PCR-based gene disruptions in Saccharomyces cerevisiae. Yeast 10: 1793–1808.
28. Gietz RD, Schiestl RH (2007) High-efficiency yeast transformation using the LiAc/SS carrier DNA/PFG method. Nat Protoc 2: 31–34.
29. Kahn M, Hasenbrink G, Lichtenberg-Frate H, Ludwig J, Kuchichino M (2010) Genetic diversity and breeding studies. J Mammal 57: 1–18.
30. Pal C, Hurst LD (2003) Evidence for inversion polymorphism related to sympatric host race formation in the apple maggot fly, Rhagoletis pomonella. Genetics 163: 939–953.
31. Gietz RD, Schiestl RH (2007) High-efficiency yeast transformation using the LiAc/SS carrier DNA/PFG method. Nat Protoc 2: 31–34.
32. Kahn M, Hasenbrink G, Lichtenberg-Frate H, Ludwig J, Kuchichino M (2010) Genetic diversity and breeding studies. J Mammal 57: 1–18.
33. Pal C, Hurst LD (2003) Evidence for inversion polymorphism related to sympatric host race formation in the apple maggot fly, Rhagoletis pomonella. Genetics 163: 939–953.
34. Wach A, Brachet A, Pohlmann R, Möller E (1994) New modular tools for PCR-based gene disruptions in Saccharomyces cerevisiae. Yeast 10: 1793–1808.
35. Gietz RD, Schiestl RH (2007) High-efficiency yeast transformation using the LiAc/SS carrier DNA/PFG method. Nat Protoc 2: 31–34.
36. Kahn M, Hasenbrink G, Lichtenberg-Frate H, Ludwig J, Kuchichino M (2010) Genetic diversity and breeding studies. J Mammal 57: 1–18.
37. Pal C, Hurst LD (2003) Evidence for inversion polymorphism related to sympatric host race formation in the apple maggot fly, Rhagoletis pomonella. Genetics 163: 939–953.
38. Gietz RD, Schiestl RH (2007) High-efficiency yeast transformation using the LiAc/SS carrier DNA/PFG method. Nat Protoc 2: 31–34.
39. Kahn M, Hasenbrink G, Lichtenberg-Frate H, Ludwig J, Kuchichino M (2010) Genetic diversity and breeding studies. J Mammal 57: 1–18.
40. Pal C, Hurst LD (2003) Evidence for inversion polymorphism related to sympatric host race formation in the apple maggot fly, Rhagoletis pomonella. Genetics 163: 939–953.
41. Gietz RD, Schiestl RH (2007) High-efficiency yeast transformation using the LiAc/SS carrier DNA/PFG method. Nat Protoc 2: 31–34.
42. Kahn M, Hasenbrink G, Lichtenberg-Frate H, Ludwig J, Kuchichino M (2010) Genetic diversity and breeding studies. J Mammal 57: 1–18.
43. Gietz RD, Schiestl RH (2007) High-efficiency yeast transformation using the LiAc/SS carrier DNA/PFG method. Nat Protoc 2: 31–34.
44. Kahn M, Hasenbrink G, Lichtenberg-Frate H, Ludwig J, Kuchichino M (2010) Genetic diversity and breeding studies. J Mammal 57: 1–18.
45. Gietz RD, Schiestl RH (2007) High-efficiency yeast transformation using the LiAc/SS carrier DNA/PFG method. Nat Protoc 2: 31–34.