Exploring and dissecting genome-wide gene expression responses of Penicillium chrysogenum to phenylacetic acid consumption and penicillinG production

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Abstract

Background: Since the discovery of the antibacterial activity of penicillin by Fleming 80 years ago, improvements of penicillin titer were essentially achieved by classical strain improvement through mutagenesis and screening. The recent sequencing of Penicillium chrysogenum strain Wisconsin1255-54 and the availability of genomics tools such as DNA-microarray offer new perspective.

Results: In studies on β-lactam production by P. chrysogenum, addition and omission of a side-chain precursor is commonly used to generate producing and non-producing scenarios. To dissect effects of penicillinG production and of its side-chain precursor phenylacetic acid (PAA), a derivative of a penicillinG high-producing strain without a functional penicillin-biosynthesis gene cluster was constructed. In glucose-limited chemostat cultures of the high-producing and cluster-free strains, PAA addition caused a small reduction of the biomass yield, consistent with PAA acting as a weak-organic-acid uncoupler. Microarray-based analysis on chemostat cultures of the high-producing and cluster-free strains, grown in the presence and absence of PAA, showed that: (i) Absence of a penicillin gene cluster resulted in transcriptional upregulation of a gene cluster putatively involved in production of the secondary metabolite aristolochene and its derivatives, (ii) The homogentisate pathway for PAA catabolism is strongly transcriptionally upregulated in PAA-supplemented cultures (iii) Several genes involved in nitrogen and sulfur metabolism were transcriptionally upregulated under penicillinG producing conditions only, suggesting a drain of amino-acid precursor pools. Furthermore, the number of candidate genes for penicillin transporters was strongly reduced, thus enabling a focusing of functional analysis studies.

Conclusion: This study demonstrates the usefulness of combinatorial transcriptome analysis in chemostat cultures to dissect effects of biological and process parameters on gene expression regulation. This study provides for the first time clear-cut target genes for metabolic engineering, beyond the three genes of the β-lactam pathway.
Background
Since the discovery of the production of antibiotics by the filamentous fungus *Penicillium chrysogenum* by Fleming in 1929 [1], much effort has been invested in selection and synthesis of strains with improved productivity [2,3]. This research has made decisive contributions to the successful large-scale production of β-lactam antibiotics after World War II. After isolation of the high-producing wild-type *P. chrysogenum* strain NRRL 1951 from a cantaloupe [4], random mutagenesis with irradiation or chemicals, followed by selection for superior production strains, enabled an over 1000-fold increase of penicillin productivity [5].

The penicillin biosynthesis pathway has been well characterized both genetically and biochemically. Biosynthesis starts with the condensation of the three amino acids cysteine, valine and α-aminoacidipic acid to form the tripeptide ACV. This reaction is catalyzed by the non-ribosomal peptide synthase ACVS encoded by pcbAB [6-8]. In the next step, the classic β-lactam ring structure is formed by isopenicillinN synthase (pcbC, IPNS) [9]. Iso-
penicillinN forms the branch point for the penicillins and cephalosporins. Penicillins can be easily produced from isopenicillinN by the exchange of the α-aminoacidipic acid moiety for a CoA activated side-chain, such as phenylacetic acid or phenoxyacetic acid, by acyl-CoA: isopenicillinN acyltransferase (penDE), which results in the production of penicillinG or penicillinV [6,7]. These three biosynthesis genes were shown to be physically linked in a penicillin biosynthesis gene cluster (pcbAB-pcbC-penDE) [7,10-13]. Present as a single copy in early strains, this gene cluster was shown to be present as amplified tandem repeats in later, high-producing strains [14,15]. Although increases in copy number have indeed been shown to result in improved productivity, saturation occurs at very high copy numbers [5,15], presumably due to limitations elsewhere in metabolism. In *P. chrysogenum* the biosynthesis pathway is compartmentalised. The enzymes involved in the first steps of biosynthesis, ACVS and IPNS, are localised in the cytosol, whereas the final steps, acyltransferase and the activation of the side-chain precursor by phenylacetyl-CoA ligase, take place in peroxisomes [16].

With some exceptions (e.g. clear increases in the copy numbers of penicillinG biosynthesis genes [14]), the molecular basis for high-level β-lactam production remains to be elucidated. Several cellular processes have been implicated in improved productivity, including a better utilisation of precursors, higher expression of biosynthesis genes, higher gene dosage and mutations in the regulatory genes controlling gene expression or other steps of the biosynthesis [12]. For rational and successful metabolic engineering, identification of these mutations will be of extreme benefit. With the availability of the *P. chrysogenum* genome sequence [17], it is now possible to study penicillinG production at a genome-wide scale.

PenicillinG production requires an exchange of the α-aminoacidipic acid side-chain of isopenicillinN for the side-chain precursor phenylacetic acid (PAA) [18]. PAA is a weak acid (pKₐ = 4.3) and as such is likely to be toxic to cells depending on its concentration and the culture pH. Such toxicity may involve specific inhibitory effects of PAA on key enzymes in biomass or penicillinG production or, alternatively, more general mechanisms such as weak-acid uncoupling [19,20]. PenicillinG producing *P. chrysogenum* can metabolise PAA via at least two routes: incorporation in the penicillinG molecule or catabolism via the homogentisate pathway to acetocacetate and fumarate [21-24]. Although this catabolic route has been described in *P. chrysogenum* [25,26] little is known about the pleiotropic effects of PAA on *P. chrysogenum*. Only a few studies dedicated to the uptake of PAA have been reported and these show contrasting results. Hillenga *et al.* [20] and Eriksen *et al.* [27] both suggest that in production strains, and at moderate concentrations of PAA, this side-chain precursor enters the cell by simple diffusion across the plasma membrane. In contrast, at low external PAA concentrations, a high-affinity transporter has been proposed to contribute to PAA uptake by the Wisconsin54-1255 strain [27,28].

The specific rate of β-lactam production by *P. chrysogenum* is strongly dependent on the availability of a suitable side-chain precursor. In many studies on the physiological impact of β-lactam production, a comparison is made between cultures grown in the absence and presence of a side-chain precursor (e.g. phenylacetic acid, phenoxyacetic acid or adipic acid) [19,29,30]. While this approach has contributed to our insight in the energetics and kinetics of penicillinG production, it does not allow a clear distinction between effects caused by, on one hand, β-lactam production and, on the other hand, PAA consumption and/or toxicity. A suitable experimental approach would be to compare two strains with similar backgrounds under well-defined culture conditions by using chemostat cultures, in which one strain would only be lacking the penicillin biosynthesis cluster.

Mutants of *P. chrysogenum* impaired in penicillin biosynthesis have been described [31-33]. Most of these mutants were derived by random mutagenesis from the low-producing strain Wisconsin54-1255, which contains one copy of the penicillin biosynthesis cluster. Although these mutants were very useful for studying gene expression and gene/enzyme relationships, for identification of the factors important for penicillinG production and PAA consumption a strain obtained from a high-producing strain background by targeted deletion is more beneficial. To
this end we constructed a strain in which the tandem-repeated penicillin biosynthesis cluster was specifically deleted, whereas the strain background was retained. In such a way it is possible to identify, by a combinatorial approach, those genes affected by phenylacetic acid and those specifically important for penicillinG biosynthesis.

Results and Discussion

Generation of a penicillin gene cluster-free derivative of a penicillinG high-producing Penicillium chrysogenum strain

To enable studies on the genome-wide impact of penicillinG biosynthesis in *Penicillium chrysogenum*, penicillin gene clusters (* PCB AB-PCB C-PEN DE*) were removed from the penicillinG high-producing strain *P. chrysogenum* DS17690. Since the β-lactam gene amplifications are in direct repeats of a larger amplified region on the same chromosome [14] spontaneous recombinations between different repeats can result in loss of gene cluster copies [15]. Isolates that underwent spontaneous recombinations were obtained via protoplast formation and subsequent sporulation of regenerating colonies. 27 random isolates were selected for Southern analysis to estimate the relative gene copy numbers of the penicillin biosynthesis gene * PCB C* and the single-copy *niaA* gene (Figure 1). The same strains were tested for penicillinV production in shake-flask cultures grown on mineral medium with the side-chain precursor phenoxyacetic acid. Half of the random isolates tested had both reduced * PCB C:niaA* ratio’s and reduced penicillinV titres. One selected strain was subjected to a second round of protoplast formation and screening. Again, 27 random isolates were selected and analysed (Figure 1B). Several putative single-copy penicillin gene cluster candidates (indicated by arrows in Figure 1B) were identified based on a similar * PCB C:niaA* ratio to that of the single-copy Wisconsin54-1255 strain [7,14,15,34]. Three of these strains were tested in for penicillinV production, with the parental *P. chrysogenum* DS17690 parent, the intermediate parent DS47274, the laboratory strain Wisconsin54-1255 and a non-producing isolate of the latter strain, npe10 [31], as controls. All three isolates showed a drastically reduced penicillinV titre, comparable to that of the single-copy strain Wisconsin54-1255. It was therefore assumed that these three isolates are derivatives of * P. chrysogenum* DS17690 that carry a single copy of the penicillin biosynthesis gene cluster (Figure 1C). To delete the last remaining copy of the penicillin gene cluster, a double homologous recombination strategy was applied. Sequences adjacent to the β-lactam biosynthetic gene cluster (3’ from the * PCB AB* gene and 3’ from the *PEN DE* gene) were used as homologous flanking sequences to target the * amds* selection marker to this locus and to delete the entire 17-kb region containing the cluster of β-lactam biosynthetic genes (* PCB AB-PCB C-PEN DE*) in the same integration event. Homologous integration by double crossover will generate transformants able to use acetamide as the sole nitrogen source (due to the presence of the * amds* gene) but unable to produce any penicillins. Moreover, genomic DNA of strains that have undergone a successful double crossover should not hybridise to β-lactam gene specific probes. Of 27,076 transformants tested, 22 (0.08%) gave no inhibition zone when overlaid with a penicillin sensitive *E. coli* strain and were selected for further analyses. All 22 putative mutants were negative in the diagnostic PCR for the β-lactam bi-synthetic gene *PEN DE*. Two of these strains gave no signal for * amds*, suggesting that they had spontaneously lost the marker gene. Southern analysis of the obtained *P. chrysogenum* strains was subsequently carried out. As a probe, a DNA fragment of 425-bp from the * PCB AB* 3’-flanking sequence was PCR amplified (Table 1, primers 19 and 20). All strains with intact penicillin biosynthesis gene sequences (i.e. the industrial parent strain as well as the single-copy copy strain) showed a 5.3-kb band in case of digestion with *HindIII* while putative β-lactam cluster-free DS17690 derivatives showed a 10.2-kb band (Figure 1D). To confirm their inability to biosynthesize β-lactams all 22 mutants were inoculated in liquid mineral medium with phenylacetic acid as precursor. Indeed, none of the mutants was capable of producing penicillinG (Figure 1E). It was concluded that all 22 strains are derivatives of the * P. chrysogenum* DS17690 strain in which all copies of the penicillin gene cluster have been deleted. Isolate DS0661 was used as cluster-free strain in this study.

Physiology of a penicillinG high-producing strain and a cluster-free derivative in chemostat cultures

The high-producing DS17690 strain and the cluster-free strain (DS0661) were grown in aerobic, glucose-limited chemostat cultures at a dilution rate of 0.03 h⁻¹, in the presence and absence of the penicillinG side-chain precursor PAA. For each combination of strain and PAA presence or absence, at least three independent chemostat cultures were analysed. In the absence of PAA, no penicillinG was produced by the DS17690 strain; however intermediates such as isopenicillinN were still produced [35]. Under the same conditions, the cluster-free strain did not produce any β-lactam intermediates. The small difference between the biomass yields cultures of the high-producing and cluster-free strains grown in the presence of PAA confirms an earlier report [36] that penicillinG biosynthesis imposes an energetic burden on *P. chrysogenum*. However, the PAA-induced reduction of the biomass yield of the cluster-free strain (Table 2) suggests that part of the biomass yield decrease that is observed upon induction of penicillinG production by PAA addition [36], may in fact be caused by the side-chain-precursor PAA itself, e.g. via uncoupling of the plasma membrane [19,20].

Although PAA could not be used for penicillinG production in the cluster-free strain, it was still consumed at circa 25% of the rate observed in the DS17690 strain. The PAA
Table 1: Strain construction

| ID primer | Gene   | Direction | Sequence                          |
|-----------|--------|-----------|-----------------------------------|
| 1         | pcbC   | FWD       | GAT TGG CGC TCC TCG TTC ACC       |
| 2         | pcbC   | REV       | CCA TTA TTT TTC TAG TCG ACA TGG CAT CGA TTC CCA AGG CCA ATG TCC CC |
| 5         | LF 7-kb| FWD       | GGT ACC                             |
| 6         | LF     | REV       | AGT TGA TAA TAT CAA TAT CTA AAA CTC CC |
| 7         | LF 5-kb| FWD       | GCC ATA TAC GAG CAT GGT ACC AGG GAC AGA TGC CCA TCC TTG |
| 8         | LF 3-kb| FWD       | GTA TAA AAG GGG AGG GTA CCC GGA AAG ATT TGT GGG CCT G |
| 9         | RF     | FWD       | GGA TGG AGC TGC GGC GCC CTC CGT CTG AAC TCC TTC GCC GCC ACT |
| 10        | RF 3-kb| REV       | CCG CCT TCC TCA CTA ACC GGC CGG CAG GTA CCG ATG GAC TCA GCA TTA TC |
| 11        | RF 5-kb| REV       | CTC TAG AAT GCT AGC GCC GTT CGA GGT ACC TTA TAG GAA AAA GGT AG |
| 12        | RF 7-kb| REV       | CCT TTT CGG TGA GGC GCC GCA ATC ACA GGT ACC GTT TTT GTC GTC |
| 13        | amdS   | FWD       | ATG CCT CAA TCC TGG GAA GAA CTG |
| 14        | amdS   | REV       | CTT GAC GTA GAA GAC GCC ACC GGC |
| 15        | penDE  | FWD       | GCC CGA GCA CAT ATG CTT CAC ATC TCT TGT CAA GCC |
| 16        | penDE  | REV       | ATG ACA AAC ATC TCA TCA GGG |
| 17        | niaA   | FWD       | CAC AGA GAA TGT GCC GTT TCT TGT G |
| 18        | niaA   | REV       | TCA CAT ATC CCC TAC TCC CGA GCC |
| 19        | pcbAB  | FWD       | GAA GAC GTC ATA CTT ATT CTC TG |
| 20        | pcbAB  | REV       | CGG CAT CGG ATA AAG AGA TCT GG |

Primers used in this study (restriction sites are in bold). LF and RF, Left and Right flanking region respectively, location of the flanking region relative to the amdS-cassette (see Figure 1A). Size of the flanking region in kb. ID primer, uniquely numbered identifier of the primer used.

consumption rate in the cluster-free strain corresponded quantitatively to the PAA consumption that was not incorporated into penicillinG in the high-producing strain (Table 2). As reported previously for industrial strains of *P. chrysogenum*, this could be the result of oxidation to 2-hydroxyphenylacetic acid by phenylacetate hydroxylase and subsequent catabolism via the homogenatisate pathway [25,27].

**Experimental design of transcriptome analysis and global gene expression responses to penicillinG production and PAA consumption**

Genome-wide transcriptome analysis was carried out on mycelium of both strains grown in glucose-limited chemostat cultures in the presence and absence of PAA at a dilution rate of 0.03 h⁻¹. The average coefficient of variation of the transcriptome data derived from independent triplicate cultures did not exceed 0.21 (Table 2), which is similar to the reproducibility obtained with chemostat cultures did not exceed 0.21 (Table 2), which is similar to the reproducibility obtained with chemostat cultures of the non-filamentous yeast *Saccharomyces cerevisiae* [37]. The level of the actA [38] and gdh2 transcripts, which are commonly used loading standards for Northern analysis, varied by less than 14% over the four situations tested. Chemostat experiments were designed to dissect gene expression responses to PAA and penicillinG production. Addition of PAA to growing DS17690 strain may induce two types of responses. Firstly, the presence of PAA itself may affect cellular processes outside β-lactam biosynthesis and thus affect the transcriptome of the fungus. Secondly, availability of a side-chain precursor enables the penicillinG formation, thus resulting in the induction and/or repression of the transcription of genes that are (in)directly related to β-lactam production. To dissect these two types of responses, we used the cluster-free strain DS50661 as a filter for the PAA response, as this strain neither produces penicillinG nor intermediates. Comparison of the transcript data of the two strains grown under the same conditions (DS17690+PAA versus DS50661+PAA; DS17690-PAA versus DS50661-PAA) will provide information on the effect of the removal of the penicillin cluster (and of possible unintended genetic changes resulting from the gene-cluster removal procedure) (Figure 2).

In total, four pair-wise comparisons between the two strains and the two conditions were performed (Figure 2), yielding a total of 1755 transcripts (representing 13% of the *P. chrysogenum* genome) that were differentially expressed in at least one of the comparisons based on the statistical criteria applied in this study ([fold difference] ≥ 2; false discovery rate 1%, see Methods section). The majority of the genome (8001 transcripts) did not show significant changes between the four conditions and transcript levels of 30% of the genome (3990 ORFs) was below the detection limit in all four situations (Figure 2).

The set of differentially expressed genes was distributed over 12 groups following a two-way comparison (Figure 3) [see also Additional file 1]. Groups 1 and 2 (Figure 3) contained genes whose transcript levels were consistently higher or lower in the presence of PAA, irrespective of the strain background. Similarly, groups 7 and 8 harboured genes whose transcript levels were consistently higher or lower in the DS17690 strain, irrespective of the presence...
Figure 1

**Obtaining a penicillin-biosynthesis-gene-cluster-free strain.** A: Double homologous recombination strategy to delete the final biosynthesis gene cluster. A denotes the pcbAB gene, B the pcbC and penDE genes and M the marker gene amended. B: Southern blot analysis to determine relative gene-copy number. Arrows indicate putative 'single copy' penicillin biosynthetic gene cluster candidates. Each number represents a single mutant. N, the non-producing isolate npe10; W, the lab strain Wisconsin54-1255, P, the parent strain DS47274, C, strain DS47276 with ±8 copies and wt, the high-producing strain DS17690. C: Relative penicillinV production by putative single copy isolates (DS47274; DS48081; DS48083; DS48088) in shake flasks. D: Southern blot analysis to characterize the cluster free strain. Genomic DNA digested with HindIII was probed with a 425-bp fragment targeting the 3’pcbAB flanking region, cluster containing strain exhibited a band at 5.3-kb while cluster free strain exhibited a band of 10.2-kb resulting from the deletion of the last copy of the cluster. E: Relative penicillinG production by putative zero-amplicon mutants (DS50650–DS50671) in shake flasks.
of PAA (Figure 3). Groups 5 and 6 represent genes that show a higher and a lower transcript level in the presence of PAA, but only in the DS17690 strain. These latter profiles are consistent with genes whose transcription is responsive to the production of penicillinG.

Enrichment of functional categories in these clusters was assessed according to the MIPS functional categories annotation [17,39] (Figure 3). To identify possible regulatory networks, a systematic search for possible protein-binding motifs in promoter sequences was performed on the different clusters.

**Gene expression responses to removal of the penicillin biosynthesis gene cluster**

A total number of 409 genes (117 in group 7 and 292 in group 8; Figure 3) showed significantly different transcript levels in cultures of the high-producing and cluster-free strains, irrespective of the presence of the side-chain precursor PAA. As expected, transcripts of the three biosynthesis genes, *pcbAB*, *pcbC* and *penDE* could not be detected in the cluster-free strain. As a result of the strain construction the other genes in the amplified region (Pe21g21280–Pe21g21420) are present as a single copy in the DS50661 strain. The observation that several of these genes showed reduced expression levels in the DS50661 strain would therefore be consistent with a gene-dosage effect (Figure 4). In addition, as confirmed by studies on a different industrial strain of *P. chrysogenum* [40] and on the laboratory strain Wisconsin54-1255 [34], not all genes in the amplified region are transcriptionally induced under penicillinG producing conditions. In addition to an enrichment of functional categories related to β-lactam biosynthesis (01.20; 01.20.37.05; 11; 11.05, 11.05.05; and 11.05.05.01), group 7 (increased transcript levels in the high-producing strain DS17690) was also enriched for functional categories 01.02 and 01.02.01, which are related to nitrogen and sulfur metabolism (Figure 3). Upregulation of the synthesis of (sulfur-containing) amino acids may be indicative for an increased synthesis of the amino-acid β-lactam precursors. The observation that higher transcript levels of these genes were also observed in the DS17690 strain when it was grown in the absence of the side-chain precursor PAA may indicate that even the low net production of β-lactam intermediates under these conditions has an impact on mRNA level regulation of precursor biosynthesis.

Among the 292 genes that were transcribed at higher levels in the penicillin-biosynthesis gene cluster-free strain than in the DS17690 strain (group 8) 17 genes were physically linked in the same chromosomal region (Figure 5). Although the increased transcript level of these genes was also observed in the presence of PAA, it was most pronounced in its absence. Annotation of many of these genes and their clustering suggests a role in secondary metabolite production (Figure 5) [41]. Among those 17 genes, the paralog of the aristolochene synthase gene, *Ari1*, from *Penicillium roqueforti* [42], which shares 97% identity, was identified (Pe12g06310). Synthesis of aristolochene represents the first step in the synthesis of sesquiterpenoids, which include potent anticancer compounds [43,44]. In *Penicillium roqueforti*, aristolochene is a precursor of PR toxin [45], for which the further biosynthesis pathway remains unidentified.

**Table 2: Physiological and microarray quality parameters for aerobic, glucose-limited chemostat cultures grown at a dilution rate of 0.03 h⁻¹**

|                      | DS17690 - PAA | DS17690 + PAA | DS50661 - PAA | DS50661 + PAA |
|----------------------|---------------|---------------|---------------|---------------|
| **Y_{x,s}** (g·g⁻¹) | 0.37 ± 0.01   | 0.35 ± 0.01   | 0.39 ± 0.01   | 0.36 ± 0.01   |
| **q_{pen}** (μmol·g⁻¹·h⁻¹) | 0.00 ± 0.00   | 19.81 ± 1.47  | 0.00 ± 0.00   | 0.00 ± 0.00   |
| **q_{PAA}** (μmol·g⁻¹·h⁻¹) | 0.00 ± 0.00   | 24.04 ± 2.38  | 0.00 ± 0.00   | 5.60 ± 0.38   |
| **q_{CO2}** (mmol·g⁻¹·h⁻¹) | 1.15 ± 0.08   | 1.42 ± 0.11   | 1.16 ± 0.05   | 1.44 ± 0.10   |
| **q_{O2}** (mmol·g⁻¹·h⁻¹)  | 1.19 ± 0.11   | 1.42 ± 0.17   | 1.16 ± 0.11   | 1.43 ± 0.08   |
| **Avg CVf**            | 0.21          | 0.18          | 0.17          | 0.14          |
| **Pc actAs**           | 4190 ± 170    | 3560 ± 360    | 3450 ± 390    | 3030 ± 680    |
| **Pc gdh2**            | 1240 ± 120    | 1140 ± 270    | 1020 ± 170    | 1060 ± 180    |
| **Repeats (n)**        | 3             | 4             | 3             | 3             |

*P. chrysogenum* strains DS17690 (penicillinG high producing) and DS50661 (lacking a functional penicillin gene cluster) were grown in the presence and absence of phenylacetic acid (PAA) in independent glucose-chemostat cultures at D = 0.03 h⁻¹. Results are the averages ± S.D. (σₚₓ).  

a: biomass yield on glucose (g of biomass per g of glucose consumed)  
b: biomass specific penicillinG production rate. In addition to penicillinG, intermediates and byproducts are formed, which accounts for ~6% of the consumed PAA [30] (μmol of penicillin G produced per g of dry weight biomass and per h)  
c: Biomass specific phenylacetic acid production rate (μmol of phenylacetic acid produced per g of dry weight biomass and per h)  
d: Biomass specific CO₂ production rate (μmol of CO₂ produced per g of dry weight biomass and per h)  
e: Biomass specific O₂ consumption rate (μmol of O₂ consumed per g of dry weight biomass and per h)  
f: the average coefficient of variation (standard deviation divided by the mean) for all genes except the genes with mean below 12  
g: encoding actin; average hybridization signal and standard deviation  
h: encoding glutamate dehydrogenase (NAD⁺ dependent); average hybridization signal and standard deviation
Gene expression responses to the side-chain precursor phenylacetic acid (PAA)

329 genes (groups 1 and 2, Figure 3) showed a consistently different transcript level in the presence and absence of PAA. Remarkably, genes belonging to the penicillin biosynthesis gene cluster were not differentially expressed in the presence and absence of PAA, with the pcb genes still being highly expressed in the absence of PAA. Only the gene encoding the PAA CoA ligase (phl, Pc22g14900) [46] showed a significantly higher transcript level in the presence of PAA (increase of 70 and 100% relative to cultures lacking PAA for the DS17690 and DS50661 strains, respectively). Recent functional characterization of phl revealed that upon its deletion, the PAA-CoA ligase activity decreased by only 40% [46], indicating the existence of one or more additional PAA-CoA ligases. Five genes with similarity to aryl- or fatty-acid CoA ligases were expressed to a higher level in cultures of both strains grown in the presence of PAA. The two closest phl homologues (Pc22g24780 and Pc22g20270) showed over 65% identity with Aspergillus proteins of unknown function, as well as a strong similarity with Arabidopsis thaliana 4-coumarate-CoA ligase (30%) (Figure 6). Further inspection of the predicted amino acid sequences of both these genes identified the presence of PTS1 peroxisomal targeting sequences. Based on these observations, Pc22g24780 and Pc22g20270 represent interesting candidates for further functional analysis aimed at identifying alternative PAA CoA-ligases in phl strains.

Both strains showed penicillinG independent metabolism of PAA. In the DS17690 strain, this was evident from the observation that PAA consumption exceeded penicillinG production (Table 2). Indeed, the PAA hydroxylase gene (pahA) [25] that encodes the first step of the PAA catabolism through the homogentisate pathway was highly induced in cultures grown in the presence of PAA (from +10- to over 100-fold). However, it has been reported, already early in the strain improvement lineage that a mutation (L181F) in this gene results in a dramatic reduction of enzyme activity [25]. Interestingly, a second gene (Pc16g01770), whose predicted protein sequence shared 42% identity with the pahA product also showed strongly elevated transcript levels (≥20-fold) in the presence of PAA. This second gene is 82% identical to Aspergillus nidulans PhacB [23], which encodes a 3-hydroxyphenylacetate 6-hydroxylase and 3,4-dihydroxyphenylacetate 6-hydroxylase cytochrome P450 monoxygenase capable of converting PAA into 2-hydroxyphenylacetate. This second gene may well be responsible for residual PAA catabolism in industrial strains that carry a loss-of-function mutation in pahA. Based on genome annotation and the transcript profiles of cultures grown in the presence and absence of PAA, the entire homogentisate pathway, which ultimately...
leads to the formation of fumarate and acetoacetate, could be mapped. Genes encoding a homogentisate dioxygenase (Pc12g09040); a maleylacetoacetate isomerase (Pc12g09020) and a fumarylacetoacetase (Pc12g09030) were tentatively identified, completing the identification of the metabolic pathway. In contrast to the other PAA-utilizing pathway (penicillinG synthesis) the five genes of the homogentisate pathway all showed a strong gene level upregulation in presence of PAA (ranging from +6.6-fold to +90-fold). Apparently, despite the reduction of the pathway's activity, transcriptional regulation of the homogentisate pathway is still functional in high-producing strains of P. chrysogenum. As indicated by their gene identity codes, the ORFs Pc12g09020, Pc12g09030 and Pc12g09040 form a chromosomal cluster. A similar clustering of the homogentisate pathways genes has been observed in Aspergillus nidulans [47]. An additional gene of this cluster, Pc12g09010, that shares 51% of identity with A. nidulans AN1893.3 and displays similarity with a putative transcription factor from Neosartorya fischeri, was also upregulated in presence of PAA in both DS17690 and DS50661 (+2.6-fold and +3.8 respectively). This observa-
tion makes it tempting to speculate that Pc12g09010 participates in transcriptional regulation of PAA catabolism (Figure 6). The characterisation of the genes encoding the pathway (a homogentisate dioxygenase (Pc12g09040); a maleylacetoacetate isomerase (Pc12g09020) and a fumarylacetoacetase (Pc12g09030)), the physical clustering of these genes and the presence of a co-clustered putative transcription factor represent interesting targets for metabolic engineering to eliminate residual rates of PAA consumption and to alleviate a potential protein burden [48].

Transport mechanisms for β-lactam antibiotics and side-chain precursors, both across the fungal plasma membrane and between intracellular compartments, remain incompletely understood. The functional category analysis of genes that showed an increased transcript level in cultures grown in the presence of PAA showed a clear enrichment of the transport-related functions (Figure 3). 42 genes in the functional categories 'cellular transport and transport mechanisms' and 'transport facilitation' showed a significantly increased transcript level in cultures grown with PAA in both strains (Figure 3, group 1).

The uptake of undissociated phenylacetic acid in *P. chrysogenum* has been reported to occur via passive diffusion [20]. However, by analogy to the well-studied non-filamentous yeast *Saccharomyces cerevisiae*, in which PAA is exported by the ATP-binding cassette transporter Pdr12 [49], its anion form is likely to be actively exported into the medium as a detoxification mechanism. From the available functional annotation of the *Penicillium* genome, 2 out the 42 genes (Pc22g14600 and...
Pc22g20390) display motif signatures of ABC transporters, as well as sequence similarity to the Aspergillus nidulans atrB [50] and atrD [51] proteins, respectively. Only Pc22g14600 belongs to the ABC-G transporters cluster [17] that also includes Pdr12, which makes Pc22g14600 a very attractive candidate for further characterization.

Activation of PAA and the final biosynthetic step in the penicillin biosynthesis pathway, the exchange of the aminoacidic acid side-chain for PAA, both occur in the peroxisome. This metabolic compartmentation of penicillin biosynthesis makes transport of isopenicillinN, PAA and penicillinG across the peroxisomal membrane an integral and important part of penicillin biosynthesis. Only one of the transporter genes that showed an increased transcript level in cultures grown with PAA (Pc21g09430) showed a clear link with peroxisomes. This gene shows strong similarity to the Saccharomyces cerevisiae ANTI gene that encodes a peroxisome-localised protein involved in adenine nucleotide transport, medium-chain fatty acid metabolism, and peroxisome proliferation [52].

The genes that showed a consistently lower transcript level in cultures grown in the presence of PAA (Figure 3 Group 2 (106 genes)), failed to show a clear enrichment of any
Dissection of gene expression responses to PAA and to penicillinG production

For future studies into the mechanism, compartmentation and regulation of penicillinG biosynthesis, it would be helpful to dissect transcriptional responses of side-chain precursor availability and penicillinG biosynthesis itself. The cluster-free strain described above synthesizes neither penicillinG nor any of its intermediates, even when grown in presence of PAA. Consequently, genes that show a transcriptional response to PAA that is specific for the high-producing strain DS17690 are likely to be related to penicillinG production rather than to the presence of PAA per se (Groups 5 and 6, Figure 3).

This approach yielded 220 genes (Group 5, Figure 3) whose transcript level was specifically higher in penicillinG producing cultures of the DS17690 strains and 305 genes (Group 6, Figure 3) that showed a lower transcript level in these conditions. Group 5 was enriched for genes whose annotation involved functions in nitrogen and sulfur metabolism, secondary metabolism and transport. Among the 18 genes assigned to the nitrogen and sulfur metabolism FunCAT annotation, six were sharing significant homology with dioxygenases (sulfur dioxygenase, taurine dioxygenase, 2-oxoglutarate-dependent dioxygenase, arylsulfatase, Fe(II)-dependent sulfonate alpha-

Figure 6
Detoxification of phenylacetic acid via incorporation into penicillinG or the homogentisate pathway. Those genes (putatively) related to penicillin, biosynthesis and phenylacetate catabolism are depicted. Total RNA was obtained from P. chrysogenum strains DS17690 and DS50661, grown in the presence and absence of phenylacetic acid (PAA) in independent glucose-chemostat cultures at D = 0.03 h⁻¹ and hybridized to Affymetrix GeneChip® microarrays. The color bar indicates the range of the mean normalized transcript level value per gene. αAAA: α-aminoadipate, Cys: cysteine and Val: valine.
ketoglutarate dioxygenase, Figure 7). Although these enzymes participate in the mobilisation of sulfur from alternative sources (e.g., in the absence of sulfate [53]), the analogy with the low-sulfate response as recorded in *S. cerevisiae* [54,55] cannot be extended further. For example, consistent with the excess sulfate included in the growth media, we did not observe a transcriptional upregulation of key enzymes in sulfate assimilation (e.g. genes involved in methionine biosynthesis). However, the transcript level of the sulfate transporter gene *sutB* [56] was specifically increased by 60% in the penicillinG producing cultures of the DS17690 strain. Moreover, the *sutB* transcript level was +4.4-fold higher in the DS17690 strain than in the DS50661 strain when grown in the presence of PAA. A similar transcript profile was observed for *Pc18g03480*, which has a strong similarity with high-affinity methionine permease genes. The second *P. chrysogenum* sulfate transporter gene *sutA* [56] was transcribed at very low levels under all conditions tested. Specific transcript level increase of several genes involved in sulfur and nitrogen assimilation in penicillinG producing scenarios suggests that a drain of amino acid precursors (cysteine and possibly valine) may affect intracellular pools of these amino acids. Indeed, metabolic flux analysis of *P. chrysogenum* grown under producing and non-producing conditions showed that the flux from 3-phosphoglycerate to serine and cysteine biosynthesis was 6-fold higher under penicillinG-producing conditions [30]. One gene involved in serine and cysteine synthesis (*Pc21g03190*, encoding a putative hydroxypyruvate dehydrogenase) showed an increased transcript level in penicillinG producing cultures of the DS17690 strain. This identifies the analysis, and possibly engineering, of cysteine and valine biosynthesis as relevant activities in applied research on β-lactam production.

The transporter for penicillinG in *P. chrysogenum* is still unknown. Although it cannot be excluded that the penicillin transporter is among the 700 constitutively transcribed transporters, the 36 transporters in group 5 form

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**Figure 7**

Transcript levels of putative sulfonatase genes specifically responding to penicillinG production. Group 5 containing genes exclusively responding to penicillinG biosynthesis contains six putative sulfonatases. Total RNA was obtained from *P. chrysogenum* strains DS17690 and DS50661, grown in the presence and absence of phenylacetic acid (PAA) in independent glucose-chemostat cultures at *D* = 0.03 h⁻¹ and hybridized to Affymetrix GeneChip® microarrays.
an interesting group for initial analysis. Out of the 36 transporter-encoding genes found in group 5, 18 were assigned to transport of a wide range of nitrogen sources, including urea (2 genes), allantoate (7 genes), various amino acids including lysine and methionine (8 genes) and oligopeptides (1 gene). Whereas lysine and methionine transport might be related to the sulfur status of the cells (see above) the role of the other genes is more elusive. However, two observations on transport-related genes provide interesting leads for follow-up studies. Firstly, Pc22g11250, whose transcript profile correlated perfectly with the production of penicillin G, shows strong similarity with the A. niger gene An15g07460, which encodes an oligopeptide transporter. Interestingly transport of β-lactams through human intestinal epithelium involves an oligopeptide (di- and tripeptide) transporter [57]. We are currently investigating the possibility that this transporter is involved in penicillin G export.

The second example a priori has no relationship with the penicillin synthesis; a group of 8 genes, whose products all show similarities with the yeast allantoate transporter Dal5, exhibited a significant upregulation under penicillin G producing conditions. These genes belong to a larger genome-wide family of 30 members. Although these 30 transporters share the same description "strong similarity to Dal5", they display very different expression profiles. Without functional analysis studies on these genes, any biological interpretation of this observation would remain speculative.

Along with the penicillin G synthesis, 18 genes that could be involved in secondary metabolism were also expressed to a higher level under penicillin G producing conditions. This group harbourd two genes Pc21g23730 and Pc21g20650 that exhibit strong similarities with a feruoyl-CoA synthetase from A. niger and a 4-coumarate-CoA ligase from Arabidopsis thaliana, respectively. While the transcript levels of these genes, remained lower than those of the two PAA-inducible putative aryl-CoA ligases mentioned above, this does not rule out a possible contribution of their gene products to in vivo PAA activation, which is further supported by the putative peroxisomal targeting signal that both harbour.

**Analysis of upstream regulatory sequences**

The 800 nucleotides upstream of the ATG of groups of genes that showed a similar transcriptional regulation were analysed for cis-regulatory elements. MEME analysis identified two motifs in group 1 and 1 motif in group 2 that met the statistical criteria applied (Figure 8; see Materials and Methods for details). Although these motifs had a good coverage of the genes in the group, the most strongly regulated genes, those involved in PAA catabolism, did not contain this motif in their 800 nucleotide upstream region. In addition, some of the motifs identified are shared by the different co-regulated groups and none match any of the known binding sites of the limited set of described transcription factors (Figure 8). Possibly the long history of strain improvement of these strains has resulted in a loss of conserved motifs. Nevertheless, a similar analysis performed on a set of 53 co-regulated genes responding to PAA in the lab strain Wis54-1255 and the DS17690 strain (cluster 1, described in [17]) identified the same motifs as found in the present study (motif 1 from group 1 and motif 2 from group 7). Increasing the length of the motif to be identified resulted in the same core conserved nucleotides (data not shown). In addition, as the groups of co-regulated genes are quite large, the discriminating power of MEME may not be sufficient to identify the actual conserved and regulating motifs. Testing a small subset of the genes, the 7 genes known to be directly involved in PAA catabolism, indeed resulted in different motifs that had a very high coverage over the genes tested. However, none of these motifs passed the stringent statistical criteria applied. These preliminary results suggest that motif identification in filamentous fungi may require different approaches than hitherto applied in more intensively studied microorganisms.

**Conclusion**

80 years after Fleming’s discovery of the antibacterial activity of penicillin, research on Penicillium chrysogenum has now become accessible to genomics approaches [17]. In the present study, we have integrated microarray-based transcriptome analysis with chemostat cultivation. This approach, which has already shown to be fruitful in other organisms such as Saccharomyces cerevisiae [54,55,58,59], Trichoderma reesei [60] and Escherichia coli [61,62], enables an investigation of the effect of individual culture parameters on genome-wide gene expression regulation. Reproducibility of transcript data is often cited as an additional advantage of chemostat-based microarray analysis [63]. Although steady-state chemostat cultivation of filamentous fungi is experimentally more challenging than chemostat cultivation of non-filamentous microorganisms, the excellent reproducibility of the transcript data obtained with P. chrysogenum indicates that this does not preclude accurate and reproducible chemostat-based transcriptome analysis.

In aerobic, glucose-limited chemostat cultures of S. cerevisiae, ca. 86 % of its 6400 genes [37] showed a detectable transcript level. Of the much larger genome of P. chrysogenum, cultivation under similar conditions yielded a detectable transcript for only 67 % of the genes. Furthermore, of the 1755 genes that showed a differential transcript level under at least one of the conditions, 53 % has unknown function. This percentage is similar to the percentage of unclassified proteins throughout the whole
Predominant motifs identified in 800 nucleotide upstream region

Promoter analysis of the 800 nucleotide upstream region of sets of co-regulated genes, identified by overlapping the results of the four pairwise comparisons (groups 1, 2, 5, 6, 7, 8), using MEME. Motifs with an E-value < 10⁻⁵ and without long stretches of A and T (> 40% GC content) were included in the analyses. Consensus sequences were depicted using the web based application WebLogo, version 2.8.2 [86,87]. N denotes the number of genes with the motif compared to the number of genes in the co-regulated group.

|   |   | N       | E-value     |
|---|---|---------|-------------|
| 1 |   | 50 of 223 | 1.5·10⁻²²   |
| 2 |   | 50 of 223 | 1.7·10⁻¹⁶   |
| 5 |   | 35 of 106  | 1.4·10⁻⁷    |
| 6 |   | 41 of 220  | 4.9·10⁻⁹    |
| 7 |   | 31 of 220  | 9.4·10⁻⁶    |
| 8 |   | 50 of 305  | 3.1·10⁻³⁵   |
|   |   | 22 of 117  | 9.4·10⁻¹⁰   |
|   |   | 29 of 117  | 4.0·10⁻⁷    |
|   |   | 44 of 292  | 1.4·10⁻³⁶   |

Figure 8: Predominant motifs identified in 800 nucleotide upstream region.
genome sequence [17]. These observations illustrate the formidable challenges that remain to be addressed in the functional analysis of the genomes of filamentous fungi. The identification of gene function in *P. chrysogenum* is likely to benefit tremendously from the rapid sequencing, annotation and analysis of the genomes of other filamentous fungi, such as those of *N. crassa* [64], *A. fumigatus* [65], *A. nidulans* [47], *A. oryzae* [66], *A. niger* [67,68], *T. reesei* [69]. For example, the recent characterisation of a new sulfate transporter gene, *astA*, in *A. nidulans*, homologous to the *S. cerevisiae* *Dal5* transporter [70] enabled us to tentatively interpret the involvement of a similar gene in *P. chrysogenum* as being part of a broader sulfur-related response.

Carefully designed transcriptomics experiments can help to prioritize targets for functional analysis based on at least three criteria: (i) the absence of a detectable transcript level rules out that the gene product contributes to either fitness or industrial performance under the experimental conditions, (ii) gene expression regulation can provide insight into the possible role of gene products in an experimental context, although relations between transcript profiles and contribution to fitness are not necessarily straightforward [71-73], and (iii) the availability of possible sequence-derived information on the putative catalytic, structural or regulatory role(s) of gene products that suggest a role in fitness and/or industrial performance [63]. Based on these criteria, several priority targets for future functional analysis studies have been identified in the present study (see Results and Discussion section).

The present study demonstrates how a simple combinatorial design of chemostat experiments, involving two *P. chrysogenum* strains, can be applied to dissect effects of side-chain-precursor availability and β-lactam production. While similar approaches have previously been applied to dissect effects of oxygen availability and nutrient limitation in *S. cerevisiae* [54,55,74], this is to our knowledge the first application of such an approach to a product-forming system. Our experimental design required a strain of *P. chrysogenum* that lacked a functional penicillin gene cluster. A cluster-free strain previously described [32], was derived from the low-producing Wisconsin54-1255 strain. As we sought to maximize the difference between producing and non-producing scenarios, a new cluster-free strain (DS50661) was derived from a penicillinG-high-producing strain background (DS17690). The approach presented in this paper should be applicable to the production of other secondary metabolites in systems where production can be controlled by the addition of a precursor molecule. In *P. chrysogenum*, this might for example include the production, by engineered strains, of the cephalosporin precursors 7-amino-deacetoxycephalosporanic acid (7-ADCA) [75], adipoyl-7-amino-3-carbamoyloxymethyl-3-cephem-4-carboxylic acid (ad7-ACCCA) [76] and deacetylcephalosporin C [77].

**Methods**

**Strains**

*Penicillium chrysogenum* strains DS17690 is a high producing strain and derived from the strain improvement program of DSM [29,35]. DS50661 lacks the penicillin biosynthesis cluster and was constructed from DS17690 as described in this paper.

**Preparation of protoplasts**

Preparation of *P. chrysogenum* protoplasts was performed as described by Cantoral *et al.* [78], using Glucanex (Novo Nordisk, Bagsvaerd, Denmark) instead of Novozyme as the cell wall degrading enzyme. Protoplasts were separated from the mycelium, washed and plated on mineral medium agar [79], without phenylacetic acid, but supplemented with 15 g l⁻¹ agar to solidify and 1 M sucrose for osmotic stabilization. Regenerating colonies were transferred to plates without sucrose to induce sporulation. Spores were collected, washed with 0.85 % NaCl, diluted and plated out on YEPD agar plates (10 g l⁻¹ Yeast Extract, 10 g l⁻¹ Peptone, 20 g l⁻¹ glucose and 15 g l⁻¹ agar). Isolated colonies were transferred to mineral medium agar plates serving as stock culture plates.

**Genomic DNA isolation**

To isolate genomic DNA, *P. chrysogenum* strains were grown in mineral-medium shake-flask cultures for 48 h at 25°C and 280 rpm. Cells were harvested, washed with 0.85 % NaCl and the pellet was frozen in liquid N₂. Frozen cells were grinded using a pestle and mortar, transferred to a plastic tube and an equal volume of phenol:CHCl₃:isoamylalcohol (25:24:1). Finally, DNA mixture was vortexed vigorously, centrifuged and the aqueous phase was transferred to a fresh tube. This procedure was repeated twice, each time using a fresh volume of phenol:CHCl₃:isoamylalcohol (25:24:1). Finally, DNA was isolated from the aqueous phase by ethanol precipitation according to standard procedures.

**Estimation of penicillin biosynthetic gene cluster copy numbers**

Genomic DNA (3 µg) was digested with EcoRI, separated on a 0.6% agarose gel and transferred to a nylon membrane by vacuum Southern Blotting. Fragments of the *pcbC* and *niaA* genes were used as probes. The former probe gives an indication of the copy number of penicillin biosynthetic genes and the latter probe is a single copy gene (encoding nitrite reductase), in *P. chrysogenum* DS17690. The probe sequences were amplified using gene specific primers 1–4 (Table 1) and labelled with the ECL non-radioactive hybridisation kit (Amersham, Little Chal-
Deletion of a single-copy penicillin biosynthetic gene cluster

To delete a single copy of the penicillin gene cluster that remained after spontaneous homologous recombination events (see Results section) a double homologous recombination strategy was applied. As double homologous crossover is a rare event in P. chrysogenum, three constructs were generated with 3, 5 and 7 kb homologous flanks on each side of the amdS gene respectively (Figure 1B). The oligonucleotides used are listed in Table 1. Following PCR amplification, the fragments were cloned in pCRXL (Invitrogen, Carlsbad, USA) via TOPO T/A cloning (Invitrogen). Subsequently, all three 5′ flanking sequences (3, 5 and 7 kb) were digested with Acc65I and NotI followed by ligation in pBluescript II SK+ (Invitrogen) pre-digested with Acc65I and NotI (Table 3). The resulting plasmids carrying the 5′-flanking fragments were then digested with NotI to facilitate cloning of the 3′ flanking sequences, which were pre-digested with NotI and Eco521 (Table 3). The obtained 3-, 5- and 7-kb flanking-plasmids all had a unique NotI site between the 5′ and 3′ flanking sequences, which was used to insert the amdS gene as selection marker. This was obtained by digesting pHELY-A1 [80] with NotI and isolating the 3.1-kb PgpdA-\(\alpha_{mds}\) expression cassette. The \(\alpha_{mds}\)-containing deletion fragments were isolated after digestion with KpnI and transformed to the single copy penicillin gene cluster strains. Transformants were selected for their ability to grow on acetamide-containing plates and afterwards screened for antibiotic production by replica plating colonies on mineral medium and overlaying them after 4 days of growth with the \(\beta\)-lactam sensitive indicator organism, Escherichia coli strain ES2231 [81]. Isolates that did not show an inhibition zone were analysed via colony PCR with three primer sets: \(\alpha_{mds}\), for the selection marker (primers 13 and 14); penDE, as indicator for the presence or absence of the penicillin biosynthetic gene cluster (primers 15 and 16) and \(\alpha_{niaA}\) as an internal control (primers 17 and 18; Table 1).

Finally, the complete removal of the three penicillin biosynthetic genes was confirmed by Southern blot. Genomics DNA (3 µg) was digested by HindIII, and probed with a 425-bp fragment corresponding to the pcbAB-3′ flanking region amplified using the primers 19 and 20 (Table 1).

### Table 3: Primer sets used for construction of double homologous crossover cassettes

| Flank  | Size (kb) | ID primer | Introduced restriction site | ID primer | Introduced restriction site |
|--------|-----------|-----------|------------------------------|-----------|------------------------------|
| Left   | 7         | Acc65I    |                              | 6         | NotI                        |
| Left   | 5         | Acc65I    |                              | 6         | NotI                        |
| Left   | 3         | Acc65I    |                              | 6         | NotI                        |
| Right  | 3         | NotI      |                              | 10        | Eco521, Acc65I              |
| Right  | 5         | NotI      |                              | 11        | Eco521, Acc65I              |
| Right  | 7         | NotI      |                              | 12        | Eco521, Acc65I              |

### Media and media composition

The mineral medium was prepared as described [35] and contained per litre of demineralised water 7.5 g glucose, 3.5 g (NH₄)₂SO₄, 0.8 g KH₂PO₄, 0.5 g MgSO₄·7H₂O, 10 mL of a trace element solution. The trace element solution contained 15 g·L⁻¹ Na₂EDTA·2H₂O, 0.5 g·L⁻¹ CuSO₄·5H₂O, 2 g·L⁻¹ MnSO₄·7H₂O, 2 g·L⁻¹ FeSO₄·7H₂O, and 0.5 g·L⁻¹ CaCl₂·2H₂O. Production of penicillinG was induced by adding 0.58 g·L⁻¹ phenylacetic acid (PAA) to the medium. To ensure similar residual concentrations of PAA, 0.30 g·L⁻¹ was added to the medium of the cluster-free strain DS50661.

### Chemostat cultivation

Aerobic glucose-limited chemostat cultivation was performed at 25°C in 3-litre turbine-stirred bioreactors (Applikon, Schiedam, The Netherlands) with a working volume of 1.8 L. The pH was maintained at 6.5 via automated addition of 2 M NaOH (ADI 1030 biocontroller, Applikon, Schiedam, The Netherlands). The fermentor was sparged with air at a flow rate of 0.9 L·min⁻¹ using a Brooks mass-flow controller (Brooks Instruments, Hatfield, USA) and stirred at 750 rpm. The dissolved-oxygen concentration was continuously monitored with an oxygen electrode (Applisens, Schiedam, The Netherlands). Continuous cultivation was initiated after 50–60 h of batch cultivation. The feed medium was supplied continuously by a peristaltic pump (Masterflex, Cole Parmer, USA) and the dilution rate was set at 0.03 h⁻¹ for all chemostat experiments. Effluent was removed discontinuously by means of a special overflow device, which has been described previously [82]. The time interval between effluent removals was fixed in such a way that each time approximately 1% of the culture volume was removed. To prevent excessive foaming, silicone antifoam (10% vol/vol, BDH Chemicals Ltd, Poole, UK) was discontinuously added at timed intervals. The offgas was cooled by a condensor at 4°C after drying with a Perma Pure dryer (type...
MD-110-48P-4, Perma Pure, Toms River, USA) oxygen and carbon dioxide concentrations were determined with a NGA 2000 analyser (Rosemount Analytical, Orville, USA). Off-gas flow rates were determined from an average of 10 measurements using a SAGA digital flow meter (Ion Science, Cambridge, UK). Specific rates of carbon dioxide and oxygen consumption were calculated as described previously [83].

Determination of culture dry weight
Culture samples (10 mL) were filtered over preweighed glass fiber filters (Type A/E, Pall Life Sciences, East Hills, USA). The filters were washed with demineralised water and dried for 20 min at 600 W in a microwave oven and were subsequently weighed.

Substrate and metabolite analysis
Glucose concentrations in the medium were determined by HPLC using an Aminex HPX-87H column (Biorad, Hercules, USA) at 60°C with 5 mM H2SO4 as the mobile phase. Phenylacetic acid and penicillinG concentrations were determined by isocratic HPLC using a Platinum EPS C18 column (Alltech, Deerfield, USA) at 30°C. The mobile phase consisted of 5 M acetonitrile with 5 mM KH2PO4 and 6 mM H3PO4.

Sampling and RNA extraction procedures
60 mL of culture broth was sampled and rapidly filtered over a glass fiber filter (Type A/E, Pall Life Sciences, East Hills, USA). The filter with mycelium was wrapped in aluminium foil, quenched in liquid nitrogen and stored at -80°C until further use. For total RNA extraction, half of the pellet was ground by mortar and pestle under constant cooling with liquid nitrogen. The powder was dissolved in 5 mL of Trizol reagent (Invitrogen) and 1 mL phenol/chloroform/isoamyl alcohol (5:1, pH 4.8, Sigma) and mixed well. The two phases were separated by centrifugation (4600 g, 15 min). Total RNA was isolated using a phenol/chloroform extraction method, which consisted of two extraction steps in acid-phenol/chloroform/isoamyl alcohol (5:1, pH 4.8, Ambion, Foster City, USA), followed by a chloroform extraction step. Each time the phases were separated by centrifugation (4600 g, 15 min). Total RNA was precipitated for 30 min at -20°C in 96% ethanol and 0.3 M sodium acetate. The RNA was collected by centrifugation (4600 g, 15 min) and dissolved in RNAse free H2O.

Microarray analysis: probe preparation and target hybridisation
Double stranded cDNA synthesis was carried out using 10 μg of total RNA and the components of the One Cycle cDNA Synthesis Kit (Affymetrix, Santa Clara, USA). The double-stranded cDNA was purified with the GeneChip Sample Cleanup Module (Affymetrix/Qiagen) followed by in vitro transcription and labelling using the GeneChip IVT labeling Kit (Affymetrix). Finally, labelled cRNA was purified (GeneChip Sample Cleanup Module, Affymetrix/Qiagen) prior to fragmentation. 15 μg of fragmented, biotinylated cRNA was hybridised to Affymetrix custom-made *Penicillium chrysogenum* GeneChip® microarrays (array code DSM_PENa520255F) at 45°C for 16 h as described in the Affymetrix users’ manual. Washing and staining of arrays were performed using the GeneChip® Fluidics Station 400 and scanning with the Affymetrix GeneArray Scanner 3000.

Data analysis
Acquisition and quantification of array images were performed using Affymetrix GeneChip Operating Software (GCOS version 1.2). Before comparison, all arrays were globally scaled to a target value of 100 using the average signal from all gene features. To the 15,531 transcript features on the arrays, a filter was applied to extract 13,746 open reading frames of which there were 13,485 different genes. This discrepancy was due to several genes being represented more than once. To represent the variation in triplicate measurements, the coefficient of variation (S.D. divided by the mean) was calculated. When the genes were ranked according to increasing average intensity, the average coefficient of variation showed a sharp increase for the genes with the lowest expression. Therefore, all genes in which the average expression in all conditions was below 12 were removed from the dataset. Subsequently all remaining values below 12 were set to a value of 12. To assess differential expression, the Significance Analysis of Microarrays (SAM version 1.21) add-in to Microsoft Excel was used for comparisons of replicate array experiments [37,84]. SAM identifies genes with statistically significant changes in expression by assimilating a set of gene-specific t tests. Each gene is assigned a score on the basis of its change in gene expression relative to the standard deviation of repeated measurements for that gene. Genes with scores greater than a threshold are deemed potentially significant. The percentage of such genes identified by chance is the false discovery rate (FDR). To estimate the FDR, nonsense genes are identified by analyzing permutations of the measurements. The threshold can be adjusted to identify smaller or larger sets of genes, and FDRs are calculated for each set [84]. Here internal SAM parameters for fold-change threshold and the false discovery rate values were set at 2 and 1% respectively.

The genes with significantly changed expression in one of the comparisons were arranged in groups via overlapping them in Microsoft Excel.

Enrichment of MIPS categories (version 1.3) was assessed by Fisher’s Exact test employing hypergeometric distribution with a p-value cut-off of 10−4 (with a Bonferroni cor-
lection). The probability was calculated as follows: the p-value of observing z genes, belonging to the same functional category is:

\[ P = \sum_{x=\max(N,M)}^{\min(N,M)} \left( \begin{array}{c} N \\ x \end{array} \right) \left( \begin{array}{c} G-N \\ M-x \end{array} \right) \left( \begin{array}{c} G \end{array} \right) \], where N is the total number of genes in a category, M is the total number of differentially expressed genes in the cluster and G is the total number of *P. chrysogenum* genes.

Promoter analysis was performed using the web-based software Multiple Em for Motif Elucidation (MEME) [85] incorporated in the software package Genedata Phylosopher (Genedata, Basel, Switzerland). The promoters (from -800, 0) of each set of co-regulated genes were analysed for overrepresented decanucleotides. Promoters with an E-value < 10^{-5} and without long stretches of A and T (> 40% GC content) were included in the analyses. Consensus sequences were depicted using the web based application WebLogo, version 2.8.2 [86,87]. The transcriptome data analysed in this study have been deposited at the Genome Expression Omnibus database http://www.ncbi.nlm.nih.gov/geo/ under the accession number GSE12632.

**Authors’ contributions**

DMH carried out fermentations. DNA microarray analysis and drafted the manuscript, ZAvdK carried out biochemical analysis, LMR and SH carried out strain construction, MAvdB and RALB coordinated strain construction and biochemical analysis, ITP designed the experiments and helped writing the manuscript. IMD designed the experiments, performed the data analysis, drafted the manuscript and coordinated the study. All authors read and approved the final manuscript.

**Additional material**

**Additional file 1**

Composition of the Group clusters (Group 1 to 12) as edited in figure 3A. Each sheet contains the expression data of the genes belonging to the group in the strains D517690 and D55066 grown in absence and presence of phenylacetate. The data shown correspond to the average of the triplicate measurements.

Click here for file [http://www.biomedcentral.com/content-supplementary/1471-2164-10-75-S1.xls](http://www.biomedcentral.com/content-supplementary/1471-2164-10-75-S1.xls)

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