Time Course Global Gene Expression Analysis of an In Vivo Candida Biofilm

Jeniel E. Nett, 1, 3 Alexander J. Lepak, 1 Karen Marchillo, 1 and David R. Andes 1, 2

Departments of 1 Medicine, 2 Medical Microbiology and Immunology, and 3 Cellular and Molecular Biology, University of Wisconsin, Madison, Wisconsin

Candida infection of devices is common and invariably associated with biofilm growth. Exploratory microarray studies were undertaken to identify target genes associated with biofilm formation from an in vivo catheter model over time. We compared messenger RNA levels from Candida albicans grown in an in vivo central venous catheter biofilm model at 12 h (intermediate growth) and 24 h (mature) to in vitro planktonic cells without a biofilm substrate, using C. albicans oligo arrays. A total of 124 transcripts were similarly up-regulated at the 12- and 24-h time points. Ontology categories most highly represented included energy/metabolism (12%), carbohydrate (10%), and protein (13%) synthesis and modification, and transport (6%). Numerous genes were previously identified from in vitro biofilm studies. These genes included those associated with hyphal growth, amino acid metabolism, adherence, drug resistance, ergosterol biosynthesis, and β-glucan synthesis. In the current data set, adherence genes were unique to those from the earlier time point. Differences between the current in vivo biofilm expression data and that previously reported from in vitro models, including alterations in metabolism and carbohydrate processing, may be due to the continuous availability of nutrients from host serum and the incorporation of the host-pathogen interaction.

Candida albicans is an important human pathogen causing a wide spectrum of diseases and is a leading cause of nosocomial bloodstream infection. Candidemia is frequently associated with implantation of a medical device. When Candida organisms grow on surfaces, such as a venous catheter or urinary catheter, they adapt to a biofilm lifestyle [1, 2]. Biofilm growth involves phenotypic changes distinct from planktonic growth [3–5]. Two such characteristics, drug resistance and reduced susceptibility to the host immune system, contribute to the difficulty in treating Candida biofilm infections. Greater understanding of the gene expression patterns present in Candida biofilms is one method to begin to discern the pathways important for the biofilm phenotype and identify potential antifungal targets.

Many in vitro model systems have been developed to mimic the biofilm growth found on infected medical devices. These models have provided the foundation for investigation of biofilm composition, architecture, and mechanisms of drug resistance [3, 5, 6]. More recently, transcription profiling experiments have identified potential biochemical pathways required for biofilm formation [7–9]. Although the substratum common to these clinical infection sites (i.e., catheter and denture material) can be used in vitro, the models lack other environmental conditions potentially important in human infections [10]. In device-associated infections, the substratum has been preconditioned with host proteins soon after implantation [1, 10]. This preconditioning film impacts biofilm formation and extracellular matrix production. In vitro models are not necessarily exposed to all the host proteins needed for the appropriate preconditioning film. Also, in many in vitro models, nutrients are depleted and waste products accumulate. However, in vivo biofilm cells are exposed...
to a continuous nutrient supply and waste products are eliminated while in the vasculature or other host tissue site [11]. Imaging of in vivo biofilms has also demonstrated incorporation of host cells, which may potentially impact biofilm formation. This host-pathogen interaction is not represented in these in vitro models. Finally, flow dynamics of host vasculature are difficult to reproduce in vitro.

In the current experiments, we use an in vivo rat catheter biofilm infection model [11]. This central venous catheter rat model mimics conditions encountered in human infections, including exposure to host proteins and immune components, host vasculature, and biofilm substratum. Inclusion of these variables in the model system allows us to account for the host-pathogen interaction and examine its impact on the transcriptional profile of Candida during biofilm development. The importance of the host-pathogen interaction has been previously demonstrated by use of in vivo model system in both the host and pathogen transcriptomes [12]. The current study demonstrates the relevance and importance of incorporating the host infection site and immune system for the biofilm process.

METHODS

In vivo venous catheter biofilm formation. A C. albicans central venous catheter biofilm model was used for in vivo experiments as previously described [11, 13, 14]. After catheter placement for 24 h, a C. albicans SC5314 inoculum of 10^6 cells/mL in a sodium chloride concentration of 0.15 mol/L was instilled in the catheter. Catheters were removed for RNA isolation or imaging after a 12- or 24-h incubation period. To yield a sufficient amount of RNA for the microarray analysis, Candida cells from 2–5 catheters were pooled. Collected cells were flash frozen with liquid nitrogen in AE buffer (sodium acetate concentration, 50 mmol/L; pH 5.2; EDTA concentration, 10 mmol/L) before RNA processing as previously described [11]. RNA was collected using the hot method as previously described [11]. The RNA integrity was assessed using Agilent Bioanalyzer 2100 with an RNA nanochip before use in microarrays.

Planktonic culture conditions. C. albicans SC5314 was propagated overnight in YPD medium at 37°C on an orbital shaker at 250 rpm and used for inoculation of planktonic cultures. Planktonic cultures were obtained in parallel with in vivo conditions by incubating the cells in 20 mL of RPMI in a glass flask at 37°C on an orbital shaker at 250 rpm. Log-phase planktonic cells were harvested by centrifugation, and the cells were flash frozen with liquid nitrogen in AE buffer until RNA processing. RNA was collected and its integrity investigated as described above.

Biofilm scanning electron microscopy (SEM). Catheters were harvested at each time point for imaging as previously described [11]. Samples were imaged in a JEOL JSM-6100 in the high-vacuum mode at 10 kV. The images were assembled using Adobe Photoshop 7.0.

Microarray and reverse transcription–polymerase chain reaction (RT-PCR). C. albicans oligo microarray slides were purchased from the Biotechnology Research Institute, National Research Council, Montreal (Version Oligo Trial 2). Three biological replicates and dye swaps were performed for each condition and time point as previously described [12, 15, 16]. The microarrays were scanned with an Agilent DNA microarray scanner (G2565BA). Lowess normalization and statistical analysis were performed using Genespring v.7 (Agilent Technologies). Genes whose expression was either increased 1.5-fold (log2) or decreased 0.67-fold for C. albicans biofilm samples relative to planktonic samples were considered for further analysis. A 2-sided Student t test was used determine the statistical significance of the log ratios.

Quantitative real-time RT-PCR was used to confirm messenger RNA abundances for a subset of up-regulated transcripts from the microarray studies. The TaqMan probe and primer sets were designed using Primer Express (Applied Biosystems) (table 1). The QuantiTect probe RT-PCR kit (Qiagen) was used from the microarray studies. The TaqMan probe and primer sets were designed using Primer Express (Applied Biosystems) (table 1). The QuantiTect probe RT-PCR kit (Qiagen) was used in an ABI Prism 7700 v1.7 sequence detection system (Applied Biosystems) as previously described [11]. Reactions were performed in triplicate. The quantitative data analysis was completed using the C(2) (−ΔΔCt) method [17]. The comparative expression method generated data as transcript fold-change normalized to a constitutive reference gene transcript (ACT1) and relative to a control or baseline condition. The statistical significance of differences in expression between biofilm and planktonic cells was analyzed by analysis of variance.

RESULTS

In vivo Candida biofilm model. Images of the intermediate biofilm (12 h) demonstrated yeast and hyphal cells attached to the substratum. At 24 h, the mature biofilm consisted of an extensive network of yeast and hyphae embedded in strands of heterogeneous matrix material (figure 1). Host cells, including red and white blood cells, were also embedded in the Candida biofilm.

Transcript profiles of intermediate and mature in vivo Candida biofilms. The genome-wide changes in gene expression...
in intermediate and mature Candida biofilms are shown in tables 1 and 2. From the intermediate biofilm and planktonic comparison, we identified 545 transcripts (8.6%) of the 6354 ORFs as differentially regulated at least 1.5-fold (table 2). Within this group, 457 transcripts were up-regulated during biofilm formation (biofilm:planktonic ratio, >1.5) and 88 transcripts were down-regulated (biofilm:planktonic ratio, <0.67). In the comparison of mature (24 h) biofilm cells and planktonic cells, 1034 transcripts (16.3%) were differentially regulated (table 2). A total of 523 transcripts were up-regulated during mature biofilm formation, and 511 transcripts were down-regulated.

Of the genes differentially expressed in intermediate biofilm formation, multiple functional categories were represented (table 2). During intermediate biofilm development, transcripts involved in protein synthesis (17%), transport (5%), stress (4%), amino acid metabolism (4%), cell wall metabolism (3%), energy and general metabolism (9%), and carbohydrate processing (6%) were abundant. The functions of 44% of the genes with transcriptional up-regulation in intermediate biofilm formation have not yet been identified. Transcripts reduced during intermediate biofilm formation were primarily involved with DNA processing and cell cycle (7%), although changes in transcription and protein synthesis (10%), transport (5%), cytoskeleton (5%), and carbohydrate processing (6%) were also evident.

In the mature biofilm, we noted similar changes in gene regulation (table 2). Again, multiple functional categories were represented. Transcription of genes associated with protein processing (17%), energy and metabolism (8%), transport (5%), DNA/cell cycle (5%), and carbohydrate processing (5%) was up-regulated in the mature biofilm cells. Similar to the intermediate biofilm, mature biofilm formation was associated with reduction of transcripts involved in DNA processing and cell cycle (7%), transcription and protein synthesis (13%), and transport (7%). The functions of nearly 40% of the differentially regulated genes are not yet known.

Next, we identified transcript abundance in both the intermediate biofilm and mature biofilm cells as a marker for relevance throughout the majority of biofilm formation. Transcript abundance of 124 genes was found at both the 12- and 24-h time points, compared with cells growing in planktonic conditions (table 3). Genes involved in carbohydrate synthesis and processing (10%), transcription and protein synthesis (13%), and energy and metabolism (12%) were among those highly expressed at both time points. Transcripts involved in cell wall metabolism (4%), iron metabolism (3%) and amino acid metabolism (5%) were also abundant.
only 27 genes was down-regulated in both the intermediate and mature biofilms (table 3). Genes involved in DNA processing (30%) were most represented among the reduced transcripts.

**Validation of microarray data by real-time RT-PCR.**
Thirteen genes of interest were chosen to include several functional categories, including those involved in adherence, drug resistance, ergosterol synthesis, transport, amino acid metabolism, cell wall synthesis, and iron metabolism. We found that the expression direction for each gene chosen was consistent with the array data (table 4).

**DISCUSSION**

Nearly all device-associated infections are associated with organisms growing as biofilms [1, 2, 18]. Most of our knowledge about *Candida* biofilms is based on in vitro studies, but such systems cannot completely simulate the environment of the infection site and host immune system. The in vivo central venous catheter model was developed to incorporate these experimental variables and has been shown to be useful for the investigation of *Candida* biofilm development, diagnosis, drug treatment and resistance, and study of the role of specific individual gene products [11, 13, 19, 20]. The current investigation expands the examination of the molecular basis for biofilm formation by assessing the temporal global transcript profile during *C. albicans* biofilm development.

These experiments identified differential regulation of many genes previously shown important in biofilm formation in vitro [4, 21–23]. Also, a subset of differentially expressed transcripts in the current in vivo study has been previously identified in genomic investigation of the planktonic *Candida* response to the mammalian host and specific host cells [12, 24]. Among the similarities to prior biofilm study was the up-regulation of genes involved in adherence (*ALS1* and *ALS4*), cell wall metabolism (*ECE1* and *SAP5*), carbohydrate and general metabolism (*CGT, ICL1, MLS1, PCK1,* and *PDK1*), and hyphal formation (*HWP2*).

Noteworthy, in vivo biofilm formation involved differential regulation of the glyoxylate cycle, which allows organisms to use 2 carbon molecules as an energy source. Transcripts for 2 enzymes of the glyoxylate cycle, *MSL1* and *ICL1*, encoding malate synthase and isocitrate lyase, were 2.4- and 2.3-fold more abundant in the mature biofilm. Increased expression of these genes has been described upon inoculation and infection of animal models in vivo and by phagocytosis [24, 25]. Mature biofilms are heterogeneous, with cells presumably exposed to gradients of nutrients. Biofilm cells closest to the catheter surface may experience a decreased supply of glucose. Increased expression of the glyoxylate cycle potentially allows the cells to use additional carbon sources.

The current analysis identified several expression differences between the 12- and 24-h biofilm time points. These differences may be accounted for by the stepwise morphologic and architectural changes during biofilm development or perhaps by a change in metabolic state or a quorum sensing process. Initial biofilm formation requires the attachment of yeast cells to a substratum [26, 27]. As expected, expression of a greater number and intensity of adherence genes was observed at the earlier 12-h time point. *ALS* (agglutinin-like sequence) genes are a family of adhesins recognized to play a role in adherence and early biofilm formation [21, 28, 29]. In the current studies, transcript abundance of *ALSI* (12-fold) and *ALS2* (4-fold) was present only at the earlier time point. The current study did not observe a transcript abundance of *ALS3*. This result supports the hypothesis that adhesins may have overlapping functions in biofilm formation in vivo. Another possibility is that *ALS3* may have been up-regulated at a time point other than those examined in the current microarray analysis.

Upon initial cell adherence to a solid surface and the initiation of biofilm formation, genes involved in amino acid biosynthesis have been shown to be differentially regulated in vitro [7, 30]. Pathways involved in sulfur metabolism and regulation of methionine and cysteine biosynthesis are among those up-regulated as early as 30 min after adhesion. In the current study, increased expression of *MET3* (1.6-fold), *MET10* (1.7-fold), *CYS3* (3-fold), and *CYS4* (1.7-fold) was detected in the intermediate biofilm. The specific role of these pathways in early biofilm formation has not yet been elucidated. *MET3* is a primary activator of sulfur assimilation in planktonic cells and is generally repressed in the presence of extracellular cysteine and methionine. However, transcript abundance of *MET3* has been identified in biofilm systems containing methionine and cysteine concentrations greater than that normally needed for repression [30]. Although it is possible that amino acids are limited within microenvironments in a heterogeneous biofilm, the swift up-regulation of these pathways upon adherence to a solid surface suggests a role in sensing and responding to the surface [30]. In contrast to the intermediate biofilm, changes in expression of genes involved in methionine and cysteine biosynthesis were not predominant in the mature phase of biofilm growth. Instead, abundance of transcripts encoding amino acid

---

**Table 2. Frequency distribution of differentially expressed genes among functional categories.**

This table is available online in its entirety at: https://www2.medicine.wisc.edu/home/infectiousdisease/timecoursesupplemental.
permeases (DAO2, DIP5, GAP6, and GNP1) was noted [31, 32].

Mature Candida biofilms are marked by a basal layer of yeast cells with subsequent layering of filamentous morphotypes and extensive matrix production [33]. Transition to the hyphal morphology intuitively requires differential regulation of hyphal-associated genes [20, 34]. The current studies identified expression of a number of hyphal-associated genes in both the intermediate biofilm (15 genes) and the mature biofilm (40 genes). Furthermore, transcription of TEC1, a transcription factor involved in hyphal morphogenesis, was up-regulated (2.4-fold) in the mature biofilm [35]. Of note, we also identified down-regulation (2.5-fold) of TUP1, a negative regulator of hyphal formation [9, 36].

Recent studies have described quorum sensing molecules in Candida organisms similar to those described in bacterial biofilm systems [37, 38]. Farnesol acts to inhibit formation of biofilm, whereas tyrosol induces replication and hyphal morphogenesis at low cell densities [39, 40]. CHK1, a histidine kinase gene, appears to be a component of the farnesol quorum sensing pathway because the chk1/chk1 mutant is unresponsive to the action of farnesol [41]. In the current study, the CHK1 transcript was abundant (1.8-fold) in the mature biofilm but not the intermediate biofilm. This timing is consistent with CHK1 involvement in the regulation of mature biofilms and, perhaps, not surprising, given the much greater burden of organisms in the milieu at this later phase of development. A recent microarray analysis compared the transcriptome of a 24-h mature biofilm exposed to farnesol with an unexposed biofilm and identified 274 differentially regulated transcripts [9]. Genes involved in hyphal morphology, drug resistance, and cell wall maintenance were highly represented. Sixty-nine of these farnesol-responsive genes were differentially regulated in the mature biofilm of the current study (table 3) [9].

We also identified altered regulation of ergosterol and β-glucan pathways associated with in vivo biofilm growth. Similar changes in gene regulation have been recently described in the most basal layer of in vitro biofilm cells, blastospores [23]. When compared with the current studies, both identified increased transcripts of ERG25 and β-1,6 glucan synthesis genes KRE1 and SKN1 [25]. ERG25 is putative C-4 methyl sterol oxidase with a role in C4-demethylation of ergosterol biosynthesis intermediates. It has been proposed that this up-regulation may allow for increased conversion of lanosterol to nonergosterol intermediates, including eburicol and 14-methyl fecosterol, at the expense of conversion to ergosterol [23, 42].

Additionally, altered expression of the β-1,3 glucan synthesis and modification pathways (FKS1, BGL2, XOG1, PHR2, FEN1, GDB1, and SGA1) was prominent in the current study [43, 44]. It has been hypothesized that the glucan pathway may restructure the biofilm cell wall and contribute to the drug-resistant phenotype [23]. The β-glucan pathway may also play a role in biofilm matrix production. Formation of a mature biofilm requires production of an extracellular polymeric matrix, which is composed primarily of carbohydrate, glucose, and protein with smaller amounts of hexosamine and phosphorus [3, 18]. The presence of glucan molecules in secreted Candida biofilm material and their possible role in biofilm resistance has recently been reported [45]. A number of transcripts involved in carbohydrate processing and synthesis [44] and cell wall metabolism [40] were differentially expressed in the mature biofilm. These gene products may participate in production of matrix material. Interestingly, transcripts involved in the β-1,3 glucan degradation pathway (ENG1 and SWC1) were reduced at the 24-h time point. Both gene products, ENG1 and SWC1, have β-1,3 glucosidase activity. Disruption of CaENG1 results in decreased extracellular β-1,3 glucanase activity [8]. It is possible that down-regulation of these β-1,3 glucosidases and altered regulation of the β-1,3 glucan pathway may serve to conserve glucans for construction of a mature biofilm matrix.

Many factors have been proposed to contribute to Candida biofilm drug resistance, including up-regulation of efflux pumps, decreased perfusion of antimicrobials through the matrix, slow growth, and alterations in plasma membrane ergosterol content [46, 47]. Early biofilm resistance coincides with increased transcript levels of the efflux pump genes MDRI and CDR1 [48]. The current studies identified transcript up-regulation of CDR2 at 12 h (1.5-fold) and MDRI at both 12 h (2.1-fold) and 24 h (1.9-fold). PDR16 transcript abundance was also noted in both the intermediate biofilm (2.2-fold) and mature biofilm (3.7-fold). PDR16 encodes a phosphatidylinositol trans-

Table 3. Transcripts differentially regulated in both intermediate and mature biofilms.

This table is available online in its entirety at:
https://www2.medicine.wisc.edu/home/infectiousdisease/timecoursesupplemental.

Table 4. Comparison of RT-PCR and microarray results for genes of interest.

This table is available online in its entirety at:
https://www2.medicine.wisc.edu/home/infectiousdisease/timecoursesupplemental.
fer protein of the Sec14p family and is up-regulated in fluco-
conazole-resistant cells overexpressing CDR1 and CDR2 [49].

Three prior studies have examined the global transcriptional response during *Candida* biofilm development [7, 9, 30]. Similar to the in vitro models, the current study found differential regulation of numerous amino acid synthesis genes in both the intermediate biofilms (18 genes) and mature biofilms (21 genes), as well as transcripts involved in sulfur metabolism (Met3 and Met10) (table 3). Analysis of the in vivo data set also identified differences from the in vitro transcriptional studies. For example, transcripts of genes encoding glucose transporters were among those most abundant at both time points. Increased expression of both HGT1 (5.4-fold) and HGT2 (52-fold) was observed in intermediate biofilms. Mature biofilm formation was associated with transcriptional abundance of HGT1 (4.7-fold), HGT2 (7.6-fold), HGT14 (2.2-fold), HGT15 (5-fold), and HGT19 (4.5-fold). One possible explanation for this expression is the need for scavenging glucose for nutrient-starved biofilm cells. Another possibility is a glucose requirement for production of the carbohydrate matrix.

A sizable subset of genes was differentially regulated in both the intermediate (8.5%) and mature (16%) biofilms. Genes involved in amino acid metabolism (AGP2, ARG1, CAN1, CDG1, CPA1, CPA2, and SMM1), cell wall metabolism (BGL2 and ECE1), iron metabolism (CFL2, FRE10, ISU1, and SIT1), and drug resistance (ERG25, PDR16, and MDR1) were among those abundant during the time periods examined (table 3). We suspect these gene products may play a role in the biofilm phenotypes exhibited throughout the biofilm lifecycle, including drug resistance, cell wall changes, altered metabolism, and maintaining cell attachment.

Of note, the study did not identify differential expression of several genes of demonstrated importance in biofilm formation (ADH1, BCRI, CPH1, EFG1, MKCl, and YWPI) [19, 21, 34, 50]. The timing of expression of these genes may not correlate with the time points examined in our study. For example, BCRI and ALS3 may play a role in earlier biofilm formation. Also, the magnitude of the differences in transcript level may be below that detectable by our microarray analysis or regulation may occur at the posttranscriptional level. The absence of these genes in the current data set points to the importance of time course experimentation.

The identification of transcripts unique to the current study demonstrates the value of in vivo models that more closely mimic the disease state. The data set provides a resource for laboratories to guide directed biofilm investigations and target specific gene products.

References

23. Khot PD, Suci PA, Miller RL, Nelson RD, Tyler BJ. A small subpop-


