Microarrays and stage conversion in Toxoplasma gondii

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Microarray technology is emerging as a powerful method for the study of those changes in gene expression that occur when a parasite converts from one life cycle stage to another. The use of appropriate experimental design and data analysis methodologies is essential if this technology is to be correctly exploited. In this review, the study of stage conversion in the transformation of Toxoplasma gondii tachyzoites into bradyzoites is considered as an example of where the application of microarrays is revealing exciting new knowledge on underlying molecular mechanisms. The more general application of microarrays in parasitology is also discussed.

Two closely related, cyst-forming coccidian parasites, Toxoplasma gondii and Neospora caninum, are economically important pathogens of livestock: T. gondii causes fetal loss in sheep and N. caninum is an abortifacient in cattle. Toxoplasma gondii is also known to occur abundantly in humans, in whom it can cause life-threatening disease in the fetus and in immunosuppressed individuals. The life cycles of the two taxa are similar, involving alternating modes of infection of the intermediate host (such as sheep or cows) or the definitive host (cats for T. gondii). However, one of the most distinguishing features of their life cycles is their persistence in tissues of an intermediate host as cystic stages, which is thus one of the reasons for their classification in the Sarcocystidae.

Tissue cysts are typically found in the central nervous system (CNS) or musculature, where they can persist for long periods of time, leading to long-term infection. Reactivation of cystic stages of T. gondii in an immunocompromised individual can result in clinical disease such as cerebral encephalitis [1], whereas recrudescence of N. caninum is thought to be associated with cases of sporadic abortion in cattle [2]. Understanding the mechanisms leading to cyst formation (cystogenesis) and their reactivation is therefore conceptually important.

Stage conversion and microarrays

Cyst formation requires the transformation of the rapidly growing tachyzoite form of the parasite to the comparatively dormant bradyzoite form. Such stage conversion in T. gondii has been studied for some years using either in vivo-generated cysts or in vitro-generated bradyzoites. Tachyzoites can be easily cultured through infection of a host cell and will differentiate into bradyzoites, and eventually cysts, under a variety of conditions in vitro (e.g. high culture pH [2,3], heat shock [2], intracellular nitric oxide (NO) [4,5], respiratory pathway inhibitors or chemicals [4,6]). In vivo, cyst formation occurs in response to interferon (IFN)-γ-mediated immunity [7]. The central dogma that has evolved from these studies is that a reduction in growth rate of the parasite results (later or simultaneously) in a switch from tachyzoite to bradyzoite, and hence the promotion of cystogenesis. This process (Figure 1) is associated with the disappearance of tachyzoite-associated markers (such as SAG1) [8,9] and the appearance of bradyzoite and cyst-related markers such as BAG1, MAG1, SAG4 and CST1 [10,11]. A great deal has been written on this specific topic [11–14], which will not be reiterated here; however, it is sufficient to point out that the temporal expression of these bradyzoite-expressed genes appears similar in all in vitro methods used to investigate stage conversion at the molecular level.

Microarrays are being extensively used now in parasitology to profile gene expression of taxa on a genome-wide scale (Box 1). The general principles behind the use of microarrays, and their strengths and weaknesses, have been described in these pages previously [15], although

Figure 1. In vitro stage conversion from tachyzoites to bradyzoites. Host cells (such as Vero) are infected with tachyzoites and cultured under conditions that promote stage conversion into bradyzoites and intracellular cysts. The transformation process is followed using a double-immunofluorescent staining method with antibodies to: (a) SAG1 (stained red); or (b) BAG1 (stained green). In (c), the specimen is stained with both antibodies and viewed under ultraviolet light. This figure was kindly provided by A. Hemphill and N. Vonlaufen (Berne). Scale (a–c) = 20 μm.
interested readers are also encouraged to look at introductory texts [16,17]. Cummings and Relman [18] provide one of the first, intriguing insights into how microarrays can be used to study microbes such as parasites, and the interactions between them and their hosts. In the first instance, microarray technology can be used simply to investigate changes in gene expression between two different cell populations, such as occurs during the conversion of tachyzoites to bradyzoites. However, it is also quite useful to try and identify groups of genes that have similar patterns of expression. Determining when and where genes are expressed allows one to make inferences about their function. For example, it has been recognized for some time that induction of bradyzoite formation in vitro results in the coordinated expression of bradyzoite-specific markers, along with the associated loss of tachyzoite-specific markers [10]. These observations have led, in part, to the raising of hypotheses that genes expressed early in the stage conversion will affect the differentiation pathway, whereas genes expressed later in the process might only affect bradyzoite maintenance and persistence. The application of microarray technology for investigating bradyzoite formation seeks to elucidate the identity and function of these intracellular processes.

Arguably, one of the greatest resources developed in parasitology over the past 10 years has been the establishment of databases containing sequence data derived from cDNAs of numerous parasites and their different life cycle stages. Large-scale sequencing of tachyzoite and bradyzoite expressed sequence tags (ESTs) have also been carried out for T. gondii [19,20] (and to a lesser extent for tachyzoites from N. caninum [21,22]), and these cDNA collections have now facilitated the production of...
microarrays that are being used for profiling changes in gene expression during stage conversion of tachyzoites to bradyzoites. Database searching and clustering of the sequence data has allowed redundancy to be limited in the microarray design, thereby maximizing the number of genes that can be included in the experimental design. A second generation of microarrays based on gene-specific oligonucleotides will undoubtedly appear.

The hope of current research, based on significant levels of investment, is clearly to understand the molecular events that occur during stage conversion and, in the longer term, this might facilitate the production of technologies that can prevent this conversion from occurring. The development of a rapid method for the generation of tachyzoites from *T. gondii* or *N. caninum* that do not encyst would clearly be of significant commercial interest. A live vaccine (using the novel S48 strain) against Toxoplasma-associated abortion in sheep is available in selected countries of the world, showing the feasibility of marketing such a product that does not persist in animal tissues [23].

**Experimental design and data analysis**

Typically in microarray analyses, one is trying to identify genes that have statistically significant changes in the level of their expression under the experimental conditions employed. Morrison and Ellis [24] provide a detailed discussion of the experimental design and data analysis techniques currently employed for this purpose (Box 2). Genes are identified that might be over- or under-expressed in the RNA populations under study, and a twofold change in expression relative to a nominated reference sample is becoming entrenched in the literature. As discussed in Ref. [24], this approach is completely arbitrary and there is no theoretical basis for choosing any ‘fold’ change as standard. Therefore, appropriate quantitative analysis of the data should involve statistical hypothesis testing in addition to recognition of patterns by multivariate analysis. Unfortunately, there is no standard protocol for performing these analyses, and so it is appropriate to try and evaluate a variety of different data analysis techniques. Nevertheless, the aims of any microarray experiment design should be to ensure the generation of high-quality data by suitably preprocessing the raw data generated, and then testing various hypotheses by statistical or multivariate analyses.

The ability to identify rigorously genes or groups of genes that are interesting in terms of their patterns of gene expression is highly dependent on adequate replication being incorporated into the experimental design, to allow for all sources of variation that might obscure the patterns one is seeking to identify (Box 3). This could occur in the form of real biological variation between experimental treatments, or through some form of variation associated with the technology (e.g. extraction of RNA, or hybridization processes). It should be recognized that each array must be considered an independent experiment, and these are the true replicates for statistical purposes. Consequently, adequate numbers of arrays should be included in the microarray experiment, preferably in a design that is statistically efficient. It is the demonstration of a common pattern(s) residing in each unique dataset. It is therefore a worthwhile strategy to examine the results of several different types of analyses, rather than merely choosing one and sticking to it – consistency of the output from a range of analyses is a good indicator that it represents a real biological pattern, whereas inconsistency is often indicative of mathematical artefacts from the particular analyses chosen. The choice of appropriate multivariate analysis techniques for microarray data is thus an area of active research.

For class comparison, several experimental groups are being directly compared. The best approach to this problem is to use traditional techniques such as analysis of variance. However, issues related to multiple hypothesis testing and probability estimation still need to be evaluated. In particular, a specific problem arises with the use of statistical tests for microarray data because a large series of statistical hypotheses are being tested (i.e. one for each of the thousands of genes) and, if a large number of statistical comparisons are made, then some of the decisions to reject the null hypothesis will be due to chance errors (false positives). Furthermore, the usual parametric assumptions that the data have a normal frequency distribution with equal variances are likely to be invalid for microarray data (i.e. the data will violate the assumptions of the analysis), and so these possibilities should not be used. Devising methods for dealing with these problems is also an active area of research.

Perhaps the most unusual aspect of data analysis for arrays is the need for normalization of the data before the analysis. Normalization is related to the technical nonreproducibility of array technology – the current manufacturing and operating characteristics are not yet good enough for unwanted sources of variation to be dealt with automatically in the laboratory protocol, and so the biologist needs to adjust the data afterwards to remove technical artefacts. Unfortunately, we do not know enough about the possible causes of these general artefacts for there to be a uniform recommended protocol that is known to be biologically appropriate. So, most of the suggestions are heuristic mathematical techniques (e.g. globalizing, smoothing, variance stabilization) that seem to work but might have as-yet-unknown less-desirable qualities. Interestingly, all of the experiments that have used vector-derived sequences for normalization, which is quite different from the more mathematical approaches usually adopted.

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**Box 2. Microarray data analysis**

Analysis of microarray data can be directed towards three distinct purposes: class comparison, class prediction or class discovery. In class comparison, data are used to test explicit hypotheses about differences between experimental groups of samples; for example, one might test whether some experimental treatment has an effect on gene expression compared with a control treatment. In class prediction, the objective is to use the data to predict the characteristics of unknown specimens based on the characteristics of known specimens; for example, an unknown DNA sequence can be assigned to one of a set of known gene classes. In class discovery, general mathematical patterns are investigated in the data, which might then be interpreted as having some biological meaning; for example, it might be possible to distinguish between different functional classes of genes based on their expression patterns.

For class prediction, a range of supervised multivariate data analysis techniques are available, and for class discovery an even broader range of unsupervised multivariate techniques have been developed. Each of these multivariate techniques has its own limitations, which need to be kept in mind when making a choice from among them. In particular, each technique searches for a different type of pattern in the data, and it might fail to find alternative but equally valid patterns. This is important because, in class prediction and discovery, the data patterns are not being specified beforehand, and it is thus necessary to choose techniques that are capable of finding the pattern(s) residing in each unique dataset. It is therefore a worthwhile strategy to examine the results of several different types of analyses, rather than merely choosing one and sticking to it – consistency of the output from a range of analyses is a good indicator that it represents a real biological pattern, whereas inconsistency is often indicative of mathematical artefacts from the particular analyses chosen. The choice of appropriate multivariate analysis techniques for microarray data is thus an area of active research.

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Irrespective of the form of data analysis, adequate replication of the arrays themselves is probably the most important current issue concerning the quantitative analysis of microarray data. Convincing evidence that a mathematical pattern is real, and not generated by random chance, can only be provided by consistency of gene expression across a series of independent experimental replicates, and the more variable the data are then the more replicates that are needed. Most microarray experiments performed to date have had inadequate replication to answer properly the experimental question being examined, and this point has been reiterated in several recent reviews.

For illustrative purposes, Ref. [24] used analysis of variance to analyze some of the data of Ref. [27], as described in the main text, and this analysis can be used to examine quantitatively how much effect each of the various components of the experiment has on the observed variation in gene expression levels. The most important of the variance components for the four-factor mixed analysis of variance are shown in part (a) of Table 1 below (there are four other components, each of which contributes < 0.1%). This analysis indicates that most of the differences in observed gene expression levels are related to the different conditions under which the cultures were grown, which is one of the two experimental manipulations. However, of more importance for our purposes here, the variation owing to differences between the replicate arrays is unduly large, larger even than the differences between the two parasite strains being studied (which is the other experimental manipulation).

From an experimentalist’s point of view, this is unacceptable because the arrays in this case are merely technical replicates, and this therefore indicates poor repeatability of the experimental manipulations. If we look at this problem in more detail, as shown in part (b) of the table, it is apparent that the experimental combination where the mutant parasite strain was grown under bradyzoite growth conditions is the culprit—the two replicate arrays for this part of the experiment are clearly not duplicates of each other. This situation is particularly unfortunate in this example, because this was the experimental combination that was predicted to have different gene expression levels compared with the other experimental combinations. The message is thus clear: array technology is currently too imprecise to be used with only a small number of technical replicates, in addition to whatever biological replicates are needed. It is therefore also important that the changes in gene expression observed in microarray experiments be confirmed by alternative methods such as reverse transcriptase (RT)-PCR.

<table>
<thead>
<tr>
<th>Component of variation</th>
<th>Contribution to total variation (%)</th>
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<tbody>
<tr>
<td>(a) Full analysis</td>
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<tr>
<td>Strains (parental versus mutant)</td>
<td>4.9</td>
</tr>
<tr>
<td>Conditions (tachyzoite versus bradyzoite)</td>
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</tr>
<tr>
<td>Expressed sequence tags (ESTs; 2066 different ESTs examined)</td>
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</tr>
<tr>
<td>Strains - conditions interaction</td>
<td>2.8</td>
</tr>
<tr>
<td>Arrays (two replicates)</td>
<td>7.2</td>
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<tr>
<td>(b) Array component only</td>
<td></td>
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<tr>
<td>Parental + tachyzoite</td>
<td>8.7</td>
</tr>
<tr>
<td>Parental + bradyzoite</td>
<td>20.1</td>
</tr>
<tr>
<td>Mutant + tachyzoite</td>
<td>2.8</td>
</tr>
<tr>
<td>Mutant + bradyzoite</td>
<td>68.3</td>
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*This example is based on the data set from Ref. [24].

**The data represent the relative percentage of how much of the total amount of variation in gene expression is ‘caused’ by each of the experimental components listed. This shows that > 7% of the variation is due simply to differences between the duplicate arrays, and that > 68% of this 7% is due to the nonreproducibility of the arrays treated with the sample from the mutant strain grown under bradyzoite conditions.”

**Microarrays for studying stage conversion**

Cleary et al. [25] reported a study on tachyzoite to bradyzoite stage conversion using microarrays constructed from ~4400 T. gondii bradyzoite cDNAs, representing at least 613 separate contigs; 95 abundant tachyzoite ESTs were included. Stage conversion from tachyzoite to bradyzoite of the BSG-4 clone of the Pru strain [26] of T. gondii was induced in vitro in human foreskin fibroblasts and changes in gene expression were investigated by comparing mRNA levels of in vitro-grown tachyzoites with mRNA levels of parallel cultures grown in high-pH media at 2, 3 and 4 days after bradyzoite induction. The expression patterns shown by the ESTs during bradyzoite induction were classified as induced, repressed, constitutive or intermediate according to the mean bradyzoite to tachyzoite transcript level determined for the EST or its contig represented on the microarray. Quantitative analyses showed that there was an inverse correlation between mRNA abundance in tachyzoites and bradyzoites for many developmentally regulated genes [25]. Independent confirmation of these relative changes in gene expression was undertaken by northern blotting. Of importance was the observation that genes previously recognized as being developmentally regulated, such as SAG1, NTP1, BAG1, SAG4A and LDH2 were all identified by microarray analyses as being differentially expressed, thereby confirming that microarray technology could faithfully display the changes in gene expression that occur during stage conversion. Most of these genes encode either surface molecules or secretory molecules from organelles associated with invasion of host cells (ROP, MIC) or intracellular development (GRA). A subset
of the genes comprised novel, differentially expressed genes that could be identified by database searching and that were not previously recognized as being developmentally regulated. The remaining contigs had no significant homology to known genes.

Cluster analysis identified several distinct classes of regulated genes that differed in their expression patterns. Many genes were not just simply switched on or off during stage conversion. For example, genes that were induced in bradyzoites fell into five categories: (i) those with transient expression (Day 2 only; Days 2 and 3; or Day 3 only); (ii) those with expression in mid-to-late stages of the conversion process (Days 3 and 4); (iii) those with expression in late stages of the conversion process (Day 4 only); (iv) those with expression at all stages (Days 2, 3 and 4); and (v) those that were all highly expressed throughout stage conversion (Days 2, 3 and 4). It was suggested that those genes that were expressed early in the stage conversion process, such as SAG2C/D and SAG4A, could be essential drivers of bradyzoite formation.

In a related study using bradyzoite-specific expression of green fluorescence protein (GFP) as a selection strategy [26], mutant T. gondii was generated by chemical mutagenesis with ethylnitrosylurea and mutants (tachyzoite to bradyzoite differentiation mutants or Tbd mutants) were identified by their reduced ability to express GFP three days after transfer to high-pH medium. These mutants, under conditions that promote stage conversion into bradyzoites, did not routinely express the CST1 cyst wall protein and had substantially reduced expression of well-recognized bradyzoite markers such as BAG1 and SAG4A. It was therefore concluded that these mutants had a global defect in the differentiation pathway leading to stage conversion. Using cDNA microarrays, the gene expression profiles of these mutants were investigated under conditions of stage conversion in vitro. Four different mutants were studied, each of which showed different characteristics in terms of their gene expression profiles. All mutants showed a decreased expression of those genes identified by Cleary et al. [25] as being induced during bradyzoite formation. Indeed, 83% of the genes that had decreased expression in the Tbd mutants are normally induced in the bradyzoite. Other categories of genes (e.g. enolase) were identified that were under-expressed in one or more mutant populations.

Other contigs were identified that are not normally induced during bradyzoite formation but were characterized by reduced expression in the Tbd mutants. These contigs were identified by their homology to 14-3-3, MAG1, ATPase A subunit, PITSLRE kinase, histone H2A variant, cyclophilin, ubiquitin-like protein and several ribosomal genes. It was hypothesized that such genes might have a regulatory role in bradyzoite formation, or might simply be dependent on common transcription factors.

Matrajt et al. [27] used insertional mutagenesis to generate stable populations of mutant parasites. A stage-specific positive/negative selectable marker based on a strong bradyzoite-specific promoter driving expression of the T. gondii hypoxanthine-xanthine-guanine phosphoribosyltransferase was used to transfect tachyzoites, and mutants in the bradyzoite differentiation pathway were identified by induction of bradyzoite formation in vitro and selection in 6-thioxanthine. One of the key advantages behind this approach is that insertional mutagenesis carried out in this way greatly facilitates the subsequent cloning and identification of the gene loci into which the insertion has occurred.

The mutant parasites grew normally even under conditions that would favor bradyzoite formation and, under culture conditions that favor stage conversion, they also showed reduced expression of markers normally associated with bradyzoites such as BAG1. Similar to the study of Singh et al. [26], RNA from a parental strain and a mutant (B7) strain grown under conditions of high pH were hybridized to bradyzoite cDNA microarrays to investigate changes in gene expression. The B7 mutant population grown under bradyzoite conditions was more similar in its patterns of gene expression to tachyzoites from either the parent or mutant strains.

Morrison and Ellis [24] illustrated the possibilities for microarray data analysis, along with flagging important experimental design issues [24]. The datasets from Refs [26] and [27] were analyzed for illustrative purposes. Although this study [24] concentrated on emphasizing the methodology of data analysis, it is interesting to make note of other important outcomes from this study. In the first instance, T. gondii mutants that failed to go through stage conversion expressed a variety of genes coding for tachyzoite-associated products such as NTPase, GRA1, GRA2, GRA5, GRA6, GRA7, undulin, toxfilin, SUL1 and ROP4. These mutants also had reduced levels of expression of the bradyzoite-specific antigens 18 and 65 kDa, SAG2- and SAG4-related mRNAs, LDH and a proteosome subunit. The main conclusion drawn from collective observations is that the four mutant strains described by Singh et al. [26] had a block in one or more regulatory pathways that prevented a commitment to bradyzoite formation from occurring, thereby giving a transcriptional profile that was ‘tachyzoite-like’. Such a conclusion was first provided by Matrajt et al. [27], as this is the most parsimonious explanation for these observations.

Second, several of the T. gondii mutants were associated with repression of a wide variety of cDNA contigs coding for ribosomal proteins such as L18, L22, L24 and S27a. The role of the ribosome in stage conversion and regulation of gene expression through translational control has largely been overlooked, even though stage conversion clearly involves clear changes in cell shape and composition which, in turn, result from changes in RNA and protein synthesis. A survey of the literature on ribosomal proteins of parasitic organisms shows that many such proteins are differentially expressed during the life cycle of a parasite. For example, in Eimeria tenella, the mRNA coding for the S3a ribosomal protein is poorly expressed in the first-generation schizont, yet strongly upregulated in the second-generation merozoite [28]; the Theileria annulata S17 protein is differentially expressed between the macroschizont and piroplasm stages [29], and components of the translational machinery, including ribosome-associated genes, are also recognized as being
differentially expressed during development of *Plasmodium falciparum* blood stages [30]. Although such evidence suggests that translational control of gene expression might be important in parasitic organisms, harder evidence in support of this conclusion is needed for *T. gondii*. Yang and Parmley [31] demonstrated that lactate dehydrogenase (LDH) is developmentally regulated during stage conversion of *T. gondii*, and that expression of LDH1 (one of the two LDH isoforms) was under translational control as mRNA coding for it could be detected in the absence of the protein [31]. Translational control also occurs during the expression of the BRS4/P36 bradyzoite-specific surface antigen (discussed in Ref. [32]). Recently, it was shown that stage conversion was associated with a downregulation of translation initiation factor 4A expression, showing that bradyzoite formation also involves a reduction in translation initiation activity [33].

Some ribosomal proteins also possess a secondary function within a cell, in addition to their essential role as a ribosome component. Roles for ribosomal proteins in essential cellular processes such as cell proliferation, replication, transcription, RNA processing, DNA repair, regulation of translation, regulation of development and apoptosis have been reported [cited in Refs 28, 29], thereby indicating the complexity of functions contributed to by ribosomal proteins. Such functional studies have yet to be undertaken in parasitic organisms like *T. gondii*.

Finally, it was possible to identify genes that possessed similar patterns of gene expression in the mutants generated in the study described in Ref. [27]. For example, the expression of ESTs 3906, 4131 and LDH (4054) were similarly repressed in all four mutants examined. Thus, one of the big benefits to be gained from undertaking microarray analyses is that it can assist in gene discovery processes through identifying genes that possess similar patterns of gene expression under the conditions studied.

**Stage conversion in *N. caninum***
The morphology (size and thickness of the cyst wall) of the tissue cyst stage of *N. caninum* and *T. gondii* are different, thereby allowing a provisional identification to be made, for example, during diagnostic investigations. Such phenotypic differences presumably reflect the different molecular compositions these stages have in the two taxa.

The ability of *N. caninum* tachyzoites to differentiate into bradyzoites in vitro has been recently reported [32,33]. Using the culture conditions established for *T. gondii*, cyst-like structures could be established, and reagents used to detect *T. gondii* cysts (such as antibodies to BAG1) were found to be crossreactive to these *N. caninum* cysts by immunofluorescence and western blotting [34,36]. The efficiency of cyst formation in vitro for *N. caninum* was not very high in comparison with *T. gondii*. An alternative in vitro culture method was described that uses continuous culture of *N. caninum*-infected murine epidermal keratinocytes for up to 8 days in the presence of 70 μM sodium nitroprusside (SNP) [35]. The expression of bradyzoite markers was confirmed by immunofluorescence, along with vacuolar staining with Dolichos biflorans agglutinin. SNP treatment also led to the downregulation of expression of the major tachyzoite surface antigen NcSAG1. A modified version of this protocol has been developed for Vero cells using lower concentrations of SNP, because it proved difficult to purify bradyzoites from keratinocytes [36].

The availability of sequence and microarray datasets in the public domain allows their analysis using a variety of methods, including the mining of databases for genes from other taxa that have not been previously recognized as important in stage conversion. Through analyses of *T. gondii* datasets, and subsequent mining of *N. caninum* sequence databases, it is possible to identify *N. caninum* genes that are implicated in stage conversion events in this species, thus allowing us to fast-track hypotheses on cyst formation in this taxon. It comes as no surprise that homologs of many well-known and well-characterized genes of *T. gondii* exist in *N. caninum* databases. BAG1, LDH and SAG4 are easily identifiable by database searching of databases, but only further investigations will tell if the molecular mechanisms of cyst formation are similar between the two taxa.

**Signaling**
Prior knowledge on the mechanism of tachyzoite to bradyzoite transformation and the formation of cysts had shown that the changes in cell shape and composition were associated predominantly with control of gene expression at the transcriptional level. The available evidence demonstrated, through detection of changes in mRNA level by northern blotting and/or reverse transcriptase (RT)-PCR, that stage conversion involved the up- or downregulation of gene expression. Tachyzoite-specific genes were switched off, whereas bradyzoite-specific genes were switched on. The studies outlined above using microarrays [24–27] show that the controlling mechanisms behind stage conversion are more complex.

One of the most important outcomes of the study in Ref. [26] was the proposal of a model covering the changes in gene expression that occur during stage conversion. A cascade representing a hierarchy of gene expression was inferred from the detected patterns of expression during the study. The presentation of this model provides a working hypothesis upon which to base future studies on the molecular events that govern stage conversion in *T. gondii* and presumably other related cyst-forming coccidian.

Some of the key features of this model must also take into consideration other known facts of cell signaling that have been elucidated in *T. gondii*. It is also possible to speculate, based on our knowledge of the mechanisms of cell signaling in mammalian cells, what the controlling mechanisms behind cystogenesis are likely to be. Changes in gene expression are normally mediated by transcription factors in response to a complex series of cell-signaling events, which are initiated from interactions at the cell membrane. It is highly likely that the response of the tachyzoite to the initial stimulus to enter stage conversion is mediated by enzymes that produce second messengers [e.g. diacylglycerol (DAG) and cyclic nucleotides]. Available evidence shows that bradyzoite formation in vitro is associated with a transient increase in cAMP concentrations [37].
Given that the process of stage conversion is associated with a reduction in growth rate, it is also necessary to invoke hypotheses that the molecular events of stage conversion and the control of replication during the cell cycle are linked. Knowledge on the cell cycle and its control in *T. gondii* is sparse. The tachyzoite cell cycle is known to contain G1, S and M phases, with little or no G2 phase [38]. In addition, a cdc2 kinase has been reported in *T. gondii* [39] and cdc2 was subsequently recognized as a key central controlling element in the *T. gondii* cell cycle, as is the case for most eukaryotes [40]. The activity of cdc2 kinases is normally regulated by a cyclin subunit, of which several are recognized. To date, only one has been described in *T. gondii* [41].

An essential role for signaling through protein kinases must also be recognized. Protein phosphorylation and/or dephosphorylation normally play a central role in cell differentiation and proliferation, and in the regulation of metabolic pathways in all eukaryotic cells. Protein kinases and phosphatases have been widely studied in many cell types, but little knowledge is available on their presence and function in parasitic protozoa. Several protein kinases have now been described in *T. gondii* and, although the complete repertoire has yet to be described, available evidence suggests that they play a central role in stage conversion. For example, phorbol 12-myristate 13-acetate (PMA) enhances cyst formation from bradyzoites in vitro [42]. Although agonists and inhibitors of protein kinases are generally nondiscriminatory in their site of action [43,44], PMA is known biochemically to mimic the activity of natural second messengers, such as DAG [45,46], which exert their effect through modulating protein kinase activity.

A mitogen-activated protein (MAP) kinase activity was demonstrated in *T. gondii* whose activity is stimulated by PMA treatment of tachyzoites [47]. MAP kinases, as the name suggests, are normally regulated by membrane-based signal transduction events leading to a cascade of kinase activities, typically referred to as the MAP kinase pathway. Ultimately, signaling through the MAP kinase pathway leads to the activation or inactivation of transcription factors in the nucleus, thereby leading to changes in gene expression. A 14-3-3 homolog exists in the cyst-forming coccidia, and in other eukaryotes this is believed to be involved in upstream signaling events leading to MAP kinase activation [48]. In mammalian cells, shaggy/glycogen synthase kinase-3 kinase is located downstream of the MAP kinase signaling cascade [49] and a homolog of this kinase is known to exist in *T. gondii*.

Inhibition of cell growth by quercetin also correlates with induction of cyst formation [50]. Although this phenomenon is mediated by changes in expression of heat-shock protein [51], it has been recognized for some time in eukaryotes that the primary mode of action of quercetin is probably at a protein kinase activity.

Conclusions

The cyst-forming coccidia are an important group of pathogens and, consequently, the generation of new knowledge on the molecular mechanisms of stage conversion, and specifically cyst-formation, is important. Large-scale sequencing projects of the *T. gondii* transcriptome have enabled the generation and application of microarrays for the study of stage conversion. This in turn has opened ‘new doors’ for the investigation of the mechanisms that underpin the morphological and structural changes observed during the formation of tissue cysts. Experimental design and the application of appropriate data analysis methodologies need to be carefully considered if microarrays are to be correctly exploited. Nevertheless, the wide application of these tools in parasitology represents an important advance for this discipline.

Acknowledgements

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