THE PROTISTAN ORIGINS OF MULTICELLULARITY: TIMING AND EVOLUTION OF CELL ADHESION MOLECULES

by

Susan Christina Sharpe

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I did it.

But ask the animals, and they will teach you;  
the birds of the air, and they will tell you;  
ask the plants of the earth, and they will teach you;  
and the fish of the sea will declare to you.  

–Job 12:7-8

"Just goes to show, we don’t know shit about fungi."

– Adam, Only Lovers Left Alive
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Abstract

Multicellularity has evolved many times within eukaryotes. Comparisons between various multicellular groups and between multicellular groups and their unicellular relatives help illuminate how this transition happened. I addressed two aspects of the evolution of multicellularity: the timing of the emergence of eukaryotic multicellular groups, and the evolutionary history of integrins, metazoan cell adhesion receptors.

Using phylogenomics and relaxed molecular clock dating methods (which employ paleontological calibrations), I estimated the timing of the emergence of eukaryotic multicellular groups. My results show that Metazoa, Fungi and two of the major multicellular red algal taxa first emerged during the mid-Neoproterozoic, whereas the dictyostelid aggregative slime moulds arose during the Paleozoic.

I found that the unicellular breviate Pygsuia biforma expresses both subunits of the metazoan integrin receptor, as well as several associated scaffolding proteins. In Metazoa, these proteins function in cell adhesion and signaling. To further study these proteins, I developed antibodies against the Pygsuia homologs of the integrin receptor proteins and the associated scaffolding protein talin, and demonstrated that they specifically recognize recombinant target proteins expressed in Escherichia coli. These tools will facilitate the elucidation of the role of integrins in Pygsuia, furthering our understanding of the ancestral functions of proteins associated with multicellularity.
### List of Abbreviations Used

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>AA</td>
<td>amino acid</td>
</tr>
<tr>
<td>AMIDAS</td>
<td>adjacent to MIDAS</td>
</tr>
<tr>
<td>ASRV</td>
<td>among site rate variation</td>
</tr>
<tr>
<td>BLAST</td>
<td>basic local alignment search tool</td>
</tr>
<tr>
<td>BLASTP</td>
<td>BLAST proteins</td>
</tr>
<tr>
<td>BS</td>
<td>bootstrap support</td>
</tr>
<tr>
<td>c-Src</td>
<td>proto-oncogene cellular-Src</td>
</tr>
<tr>
<td>Cal</td>
<td>calibration</td>
</tr>
<tr>
<td>CIR</td>
<td>Cox-Ingersoll-Ross</td>
</tr>
<tr>
<td>DHFR</td>
<td>dihydrofolate reductase</td>
</tr>
<tr>
<td>DNA</td>
<td>deoxyribonucleic acid</td>
</tr>
<tr>
<td>ECM</td>
<td>extracellular matrix</td>
</tr>
<tr>
<td>EGF</td>
<td>epidermal growth factor</td>
</tr>
<tr>
<td>FAK</td>
<td>focal adhesion kinase</td>
</tr>
<tr>
<td>FERM</td>
<td>band 4.1, Ezrin, radixin, moesin domain</td>
</tr>
<tr>
<td>GTR</td>
<td>general time reversible</td>
</tr>
<tr>
<td>HMM</td>
<td>hidden Markov Model</td>
</tr>
<tr>
<td>HRP</td>
<td>horseradish peroxidase</td>
</tr>
<tr>
<td>ILK</td>
<td>integrin-linked kinase</td>
</tr>
<tr>
<td>IMAC</td>
<td>integrin-mediated adhesion machinery</td>
</tr>
<tr>
<td>IPP complex</td>
<td>ILK, PINCH, paxillin complex</td>
</tr>
<tr>
<td>IPTG</td>
<td>isopropyl beta-D-1-thiogalactopyranoside</td>
</tr>
<tr>
<td>ITA</td>
<td>integrin α</td>
</tr>
<tr>
<td>ITB</td>
<td>integrin β</td>
</tr>
<tr>
<td>ITG</td>
<td>integrin</td>
</tr>
<tr>
<td>LBA</td>
<td>long-branch attraction</td>
</tr>
<tr>
<td>LECA</td>
<td>last eukaryotic common ancestor</td>
</tr>
<tr>
<td>LG</td>
<td>Le-Gascuel amino acid replacement matrix (Le and Gascuel 2008)</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Definition</td>
</tr>
<tr>
<td>--------------</td>
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<tr>
<td>LIMBS</td>
<td>ligand-induced metal ion–binding site</td>
</tr>
<tr>
<td>LogN</td>
<td>lognormal</td>
</tr>
<tr>
<td>LOX</td>
<td>lysyl oxidase</td>
</tr>
<tr>
<td>Ma</td>
<td>millions of years ago</td>
</tr>
<tr>
<td>MCMC</td>
<td>Markov Chain Monte Carlo</td>
</tr>
<tr>
<td>MIDAS</td>
<td>metal ion-dependent adhesion site</td>
</tr>
<tr>
<td>ML</td>
<td>maximum likelihood</td>
</tr>
<tr>
<td>MRCA</td>
<td>most recent common ancestor</td>
</tr>
<tr>
<td>mRNA</td>
<td>messenger RNA</td>
</tr>
<tr>
<td>MRO</td>
<td>mitochondrion-related organelle</td>
</tr>
<tr>
<td>Myr</td>
<td>millions of years</td>
</tr>
<tr>
<td>PCD</td>
<td>programmed cell death</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
</tr>
<tr>
<td>rDNA</td>
<td>ribosomal DNA</td>
</tr>
<tr>
<td>RGD</td>
<td>arginine – glycine – aspartate</td>
</tr>
<tr>
<td>RMC</td>
<td>relaxed molecular clock</td>
</tr>
<tr>
<td>RNA</td>
<td>ribonucleic acid</td>
</tr>
<tr>
<td>SDS-PAGE</td>
<td>sodium dodecyl sulfate-polyacrylamide gel electrophoresis</td>
</tr>
<tr>
<td>SSU</td>
<td>small subunit</td>
</tr>
<tr>
<td>TBS-T</td>
<td>tris buffered saline, 0.1% Tween 20</td>
</tr>
<tr>
<td>TS</td>
<td>thymidylate synthase</td>
</tr>
<tr>
<td>UGam</td>
<td>uncorrelated gamma</td>
</tr>
</tbody>
</table>
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Chapter 1: Introduction

One of the most interesting evolutionary questions to big, multicellular organisms such as ourselves is how unicellular organisms evolved to the complex organisms that inhabit our visible environment today. With the advent of molecular phylogenetics, the unicellular organisms most closely related to multicellular groups can be identified, and their genomes sequenced and annotated. The sequence data gathered to date reveal that many genes needed for multicellular development existed before the transition to multicellularity (Rokas 2008, Knoll 2011). Improved phylogenies of eukaryotes have led to a better understanding of the number of unicellular to multicellular transitions, while recent advances in genomics mean that the genetic signature of multicellularity is being elucidated. This chapter outlines the theory behind what is needed for the transition from unicellularity to multicellularity (both in terms of new functions required and broader selection pressures) and summarizes what is known about the origin of one multicellular group, namely animals.

1.1 Diversity of Multicellularity

The evolution of multicellularity can be seen as the result of selection for an increase in size (Bonner 2000). As surface areas grows more slowly than volume for a cell of the same shape with a growing length, surface diffusion limits how large a single cell can be (Saucedo and Edgar 2002), excluding unicellular organisms from certain niches. Cell size is related to genome size, although the mechanism of this relationship is not well understood (Cavalier-Smith 2005, Gregory 2001). Some cells have evolved new and interesting cell biology to get around these constraints, one example being the giant cells of Acetabularia, which are up to several centimeters in length (Mandoli 1998). Other large cells function by containing many nuclei (for example myxogastrids – see the ‘Multinucleate’ column in Fig 1.1). Another strategy for the organismal increase in size, multicellularity, can arise in two fundamentally different ways: either by cells aggregating or remaining attached after cell division.
Figure 1.1: Diversity of eukaryotic multicellularity. Aggregative multicellularity involves solitary vegetative cells coming together to form fruiting structures. Colonial multicellularity consists of cells remaining attached after division, while complex multicellularity involves a large enough mass of cells that some are not in contact with the outside environment (Knoll 2011). Groups straddling these columns represent intermediate grades of organization— with some cellular differentiation, but less complexity, and various forms of multicellularity within the groups. Multinucleate organization—cells that reach huge sizes with many nuclei—is an alternative strategy for the evolution of macroscopic forms. Early fungi consist of multinucleate hypha, which in some groups have developed cross-walls and a high level of differentiation in fruiting bodies (Stajich et al. 2009). Parenthesis indicate subgroups of the above group—for example kelps are a subset of brown algae and Bryopsidales are a subgroup of Chlorophyta. Pezizomycotina and Agaricomycotina are subsets of Fungi that evolved complex multicellularity independently (Stajich et al. 2009). Bold indicates possible multiple transitions to multicellularity and/or reversion to unicellularity.

Lists of colonial and multinucleate forms are not exhaustive.

Aggregative multicellularity is common throughout eukaryotes (Fig 1.1, first column). All aggregative multicellular organisms have a similar life cycle: single cells feed, then when the food runs out they come together, forming spores. In many species these spores are lifted up on a stalk, which can be made of an extracellular matrix, cells, or both (Brown and Silberman 2013). While this strategy does help the organism disperse to more favorable environments, the fact that many different cells are coming together means that, in principle, they may not all be genetically identical, leading to interesting
selection effects (discussed in more detail below). Most examples of aggregative multicellularity are simple, with only one or two different cell types. *Dictyostelium discoideum* (Amoebozoa) contains a polarized epithelium (Dickinson et al. 2011), similar to the one found in animals, making it an interesting organism for comparative studies.

Most multicellular groups, including the physically largest, develop by cells remaining attached after division (Fig 1.1, middle columns). The simplest example of this life-history is ‘colonial growth’, which simply involves cells sticking together, without any cellular differentiation. In some diatoms (for example *Skeletonema*), this is achieved by interlocking the silica shells of adjacent cells (Gebeshuber and Crawford 2006), while the rhizarian *Spongomonas* and relatives of *Volvox* extrude an extracellular matrix. Some choanoflagellates form clumps of cells (Fairclough et al. 2010) and some ciliates develop a colonial form of several cells attached to an extracellular stalk (Bonner 1998). Some colonial organisms with many cells have cells that differentiate, increasing complexity. This ranges from the simple germ/soma differentiation in *Volvox* to the more than 100 cell types in some animals (Rokas 2008). After a certain increase in organism size, differentiation becomes necessary – now some cells are separated from their environment, and will need specialized structures for the transport of nutrients (Knoll 2011).

Definitions of complex multicellularity vary, but usually rely on number of different cell types (Rokas 2008) and the presence of a recognizable macroscopic morphology (Cock et al. 2010). When defined as those organisms that are large enough that some cells are no longer in contact with the external environment (Knoll 2011), complex multicellularity has evolved in the land plants, florideophyte red algae, kelps, animals, and twice in fungi (groups entirely within the ‘complex’ column of Fig. 1.1).

Multinucleate forms are also widespread in eukaryotes (Fig 1.1, rightmost). This includes foraminiferans, the green algal group Bropsidales, some stramenopiles related to brown algae (e.g. *Vaucheria*), and the myxogastrids (or plasmodial slime molds) found in Amoebozoa. Simple, deep-branching fungi (such as the Mucoromycotina, which includes *Rhizopus*) consist of long filaments surrounded by a cell wall made of chitin, with no cell walls separating the many nuclei. In more complex forms, septa (cross walls that divide the hyphal tube into sub-compartments) separate nuclei, explaining the classification of Fungi as a multicellular organism, but even here nuclei can move
throughout the hyphae. Complex multicellularity with many cell types has evolved in both the ascomycetes and basidiomycetes (Knoll 2011), specifically the Pezizomycotina and Agaricomycotina (Stajich et al. 2009).

While the multiple distinct forms of multicellular organisms detailed above make it difficult to count the number of transitions to multicellularity, another difficulty lies in elucidating phylogenetic relationships of multicellular groups and their unicellular relatives. Early phylogenetic trees of eukaryotes placed many complex multicellular groups within the so-called ‘crown eukaryotes’ (Sogin et al. 1989), which excluded ‘simpler’ forms such as amitochondrate protists. Further study of the effects of divergent sequences on phylogenetic reconstruction revealed that this grouping of complex organisms is the result of distantly related organisms grouping together due to a long-branch attraction (LBA) artifact (Roger 1999). Theories explaining the transition from unicellularity to multicellularity rely on inferences of the ancestral unicellular state, as inferred from the extant deep-branching groups. To identify deep-branching groups, the tree must be rooted, which requires additional information, such as sequences from an outgroup, which may be divergent enough to increase the LBA artifact. Issues of eukaryotic phylogeny are discussed in more detail in the introduction to Chapter 2.

Increasing genetic distance may make the reconstruction of phylogeny difficult, and even if known unicellular species exist that are the sister-groups of multicellular taxa, their features may no longer be representative of the ancestral form. Volvox and its relatives is an example of a recent (209–260 Ma) transition that has proven very profitable to study (Herron 2009), while the transition in Metazoa is much more ancient (660 – 880 Ma; see Chapter 2 (Sharpe et al. 2015)) and accordingly more difficult to reconstruct.

While eukaryotes dominate the macroscopic biosphere, multicellularity is also present in prokaryotes (Claessen et al. 2014). Biofilms held together by secreted matrix components (for example in Bacillus subtilis) are similar to some forms of colonial development (Vlamakis et al. 2013). Other prokaryotes show differentiation, a step towards complex multicellularity. In filamentous cyanobacteria such as Anabaena, cells differentiate, losing their reproductive capacity, so that they can fix nitrogen (Golden and Yoon 2003). The regulation of this process, as well as the mechanisms for sharing the resources produced, involves complex regulation and communication between cells. In
the actinomycete *Streptomyces*, inhibition of cell division leads to the formation of hyphae – filaments with multiple nucleoids – which undergo a developmental program-to produce spores that involves programmed cell death, cooperation, differentiation and the production of antibiotics (Claessen et al. 2014). In Archaea, *Methanosarcina* grows in clumps, possibly to protect inner cells from an influx of harmful oxygen (Bonner 1998).

1.2 New Functions Required for Multicellularity

The transition from a unicellular to a multicellular lifestyle involves the acquisition of two key features: cells that previously would have existed independently must stick together, and must cooperate rather than compete. While unicellular organisms must maintain complex signalling systems to respond to their changing environments (Christensen et al. 1997), multicellularity requires communication between cells of the same organism. Signalling can help cells maintain homeostasis for the larger organism, incorporating information about the status of other cells, and control how and when cells differentiate. In some cases, cells even undergo programmed cell death if that is what is beneficial to the larger organism (Nedelcu et al. 2011). Genes that control early development are of particular interest to understand how complex body plans are formed (Richards and Degnan 2009). Adhesion and signalling proteins (and proteins that fulfill both roles) are therefore important for establishing the differentiation and cooperation needed to make a multicellular organism.

In each separate lineage where multicellularity has evolved, different ways for cells to adhere to one another have evolved (Abedin and King 2010). Plants use polymers to attach the cell wall of one cell to the next, maintaining the shape of the tissue (Jarvis et al. 2003). In contrast, animals possess a variety of cell adhesion systems, each of which fulfills slightly different roles (Abedin and King 2010). Two examples are tight junctions, which create barriers between the outside and the inside of the organism, and focal adhesions, which create connections between certain cells and the extracellular matrix. Each of these systems is made up of a specific set of proteins, and finding homologs of these proteins in unicellular organisms could shed light on what their functions were before the transition to multicellularity. Although it is not expected that unicellular
organisms would have a use for such adhesion molecules, integrins have been found in *Capsaspora owczarzaki* and *Thecamonas trahens* (Sebé-Pedrós et al. 2010, Brown et al. 2013), while cadherins have been found in choanoflagellates (King et al. 2003). Possible functions for these molecules in unicellular organisms include motility, predation or interactions during sexual reproduction.

Animals maintain the integrity of their tissues with a proteinaceous extracellular matrix (ECM) (Hynes 2012). Many extracellular matrix proteins are long, with many repeating, independently folded domains. These domains serve as sites for cross-linking with other proteins (increasing ECM stiffness), or as binding sides for transmembrane receptors, allowing cells to maintain a position in a tissue or move across it. The ECM acts as a reservoir for growth factors, regulating their accessibility and adding a spatial organization element to their binding to receptors on cells (Hynes 2009); the ECM thus has a signalling role in development. The arrangements of domains in ECM proteins are generally conserved in Metazoa, with many specific innovations in vertebrates. Many of the ECM protein domains are present in unicellular lineages such as choanoflagellates and *Capsaspora*, but with different arrangements (Williams et al. 2014). The proteins necessary for the formation of basement membrane, a structure that is present in all Metazoa except sponges, are type IV collagen, laminin, nidogen and perlecan (Hynes 2012).

One family of proteins that has only recently been recognized as existing outside of Metazoa are the lysyl oxidases (LOX). While best known for their role in modelling the extracellular matrix (by creating cross-links between collagen molecules), it was recently shown that the conserved catalytic domain of LOX has a broad phylogenetic distribution, and is found in both prokaryotes and eukaryotes (Grau-Bové et al. 2015). While the specific domain arrangements of both LOX and the proteins it modifies are confined to Metazoa, many of these domains are also found in other organisms, leaving open the possibility that their original functions were co-opted for multicellular development in the Metazoa.

The evolution of multicellularity also makes possible the simultaneous differentiation of cell types, where cells specialize for different roles, often increasing the efficiency of each of the individual roles. Differentiation may be a necessary result of
increased size, as once some cells are no longer in contact with the outside environment, the organism must contain specialized nutrient transport systems to survive (Knoll 2011). Often differentiation comes at a direct cost to individual cells: heterocysts in multicellular cyanobacteria (Rosetti et al. 2010), and the cells that form the base and stalk in dicyostelid slime molds (Thomason et al. 1999), are examples of cells that cannot reproduce after differentiating.

Many of the signalling genes important for metazoan development were present in the common ancestors of Metazoa and their unicellular relatives, but underwent domain enrichment in metazoans, perhaps concomitant with the origins of differentiated cells in this lineage. The metazoan Hippo signalling pathway, which regulates cell proliferation and apoptosis as well as organ size, has been found in the filasterean Capsaspora owczarzaki (Sebé-Pedrós et al. 2012). G-protein coupled receptors (GPCRs) are important for responding to the environment in all eukaryotes, and the cytoplasmic proteins that transduce these signals are conserved in Holozoa (Metazoans + unicellular relatives; see Fig 1.2). Metazoans show a significant expansion of the number of GPCRs, related to new functions involved in multicellularity (de Mendoza et al. 2014). Transcription factors play an important role in development in Metazoa and land plants, by changing the expression of other genes (de Mendoza et al. 2013). In particular the metazoan T-box transcription factors are present in the filasterean Capsaspora; however they have not sub-functionalized to the level present in their metazoan homologues (Sebé-Pedrós et al. 2013a).

Programmed cell death (PCD) is a mechanism in multicellular animals by which cells undergo a planned program of systematically destroying themselves. This is obviously a detrimental process for an individual cell, but can have benefits for the larger organism. Programmed cell death has evolved to prevent necrosis (which would harm other cells), and inappropriate growth (i.e.: cancer). While processes similar to programmed cell death are phylogenetically widespread across unicellular life (Nedelcu et al. 2011), their functions are less clear. While kin selection, where cooperation of closely related individuals ensures continued survival of their genes, could explain the evolution of programmed cell death in unicellular organisms, it only applies when populations are largely made up of one species, or in the presence of some mechanism for
recognizing kin (Nedelcu et al. 2011). Another option is that programmed cell death is the result of the pleiotropic effects of genes. In this scenario, genes that act in ways beneficial to the cell under some conditions may systematically destroy it under other conditions (Nedelcu et al. 2011). Selection for the genes due to their beneficial effect has the side effect of selecting for programmed cell death in some cases.

Being multicellular involves cooperating with related cells to function as an individual. Individuals must be able to distinguish self from non-self, or else they risk wasting resources on an unrelated neighbour. While less of an issue for multicellular organisms that stay together through rigid cell walls (e.g.: plants, most algae), recognition of non-self may be a greater concern for other groups such as animals, which often have cell types that move across other cells or the ECM. For organisms with dynamic cell adhesion, restricting this adhesion to ‘self’ cells (and hence being able to distinguish ‘self’ from ‘non-self’) could be an important factor in the origin of an immune system (Grice and Degnan 2015) and the maintenance of multicellular tissue against the threat of cheaters (cells that proliferate or use resources in a way that benefits the cell, but not the larger organism).

1.3 Selection Pressures that Favour Multicellularity

The evolutionary transition from unicellular to multicellular lifestyle may be the result of several different selection pressures. Increased size can be an advantage in avoidance of predation (Boraas et al. 1998), and smaller surface-to-volume ratios with increasing cell size limit the size of a single cell (Saucedo and Edgar 2002). The transition to multicellularity has been replicated in experiments, by selecting for settling through a liquid, favouring clumps of cells in yeast (Ratcliff et al. 2012) and in Chlamydomonas reinhardtii (Ratcliff et al. 2013). In another experiment, the unicellular Chlorella vulgaris transitioned to an eight-cell colonial form after exposure to a flagellate predator (Boraas et al. 1998). Another possible advantage is the improved absorption of nutrients. For yeast, more benefit is obtained from secreting enzymes that break down food if the cells are clumped together (Koschwanez et al. 2011), as when cells are in clumps, the concentration of enzymes, and by extension of available nutrients, is higher near the cell. While it may be tempting to assign selective advantages to all aspects of
multicellular growth, some features of multicellularity, such as the length of filamentous bacteria, may simply represent an emergent property of processes such as cell death and birth rates (Rossetti and Bagheri 2015).

The unicellular-to-multicellular transition is characterized by a change in the unit of selection (Grosberg and Strathmann 2007, Maynard Smith and Szathmary 1995). Organisms go from being single cells, each competing for their own survival, to a collective of cells, with tasks divided among cells in ways that increase the fitness of the group, but not necessarily the individual cells (for example, programed cell death, discussed above). For many multicellular organisms, a genetic bottleneck in one stage of the life cycle ensures that all the cells in the organism are closely related, meaning their contribution to the survival of the larger organism helps pass their genes on to the next generation (Buss 1987). While multicellular organisms have evolved mechanisms to avoid ‘cheaters’ in the collective, they are not foolproof. As a result, cancer and cancer-like phenomena are known amongst diverse lineages of multicellular organisms (Aktipis et al. 2015). The complexity of animals may leave them especially susceptible to cancer, in particular metastasis. Runaway cell growth causing tumours/galls/lesions has also been observed in fungi, plants, red algae and brown algae (Aktipis et al. 2015). Cheaters are also present in examples of aggregative multicellularity, and are well studied in dictyostelid slime moulds (Bonner 2009, Fairclough et al. 2013).

One possible explanation for the persistence of multicellular forms of life, despite their susceptibility to cheaters, is that adaptations that favour multicellularity act as a ratchet (Libby and Ratcliff 2014). Under this process, adaptations that offer selective advantages for multicellular organisms and disadvantages for unicellular organisms mean that if the organism reverts to unicellularity, it is at a disadvantage compared to other unicellular organisms. An example is found in the ‘snowflake yeast’, which evolved as a result of artificial selection for cells or clumps of cells that settle in liquid (Ratcliff et al. 2012). Increased apoptosis is an advantage for the multicellular yeast, as more clumps of several cells are released as propagules, but is a disadvantage for unicellular yeast, as more of the cells die. Once a lineage has evolved multicellularity, while it may revert to unicellularity, it is at a disadvantage compared to other unicellular lineages when it does so. The return to a successful unicellular lifestyle depends on multiple steps going against
the ‘ratchet’ of the multicellular organisation, and so is not likely to happen. Cancers can be thought of as a return to selection at the cellular level, however they can only survive within a multicellular body, and so represent an evolutionary dead end. The genetic factors that made the cancer successful within an organism make it hard for the cells to survive outside a larger body. Occasionally, cancers have become transmissible, for example is the canine transmissible venereal tumour (CTVT) (Murchison et al. 2014). This sexually transmitted disease found in dogs began as a cancer in a single dog, but was able to develop into a transmissible pathogen after many mutations.

Reconstructing the selective forces that led to extant multicellular groups can be difficult, as the details of the environment of the ancestral species are usually not known. Advantages such as avoiding predation assume the presence of larger, possibly multicellular predatory organisms, and while this may be useful to explain the origins of multicellularity in groups that have evolved relatively recently (for example, *Volvox*), it does not explain how it first happened in organisms such as Metazoa. On the other hand, the top of the size range of extant organisms represents a perpetually unfilled niche, just waiting for innovations allowing for increases in size for the niche to be exploited (Bonner 1998, Bonner 2006). Features that are necessary for the maintenance of the complex multicellular organisms we see today may have evolved under very different selection pressures.

1.4 Evolution of Life Histories and Multicellularity in Amorphea

Amorphea, the supergroup which contains both Amoebozoa and the Opisthokonta, as well as other protist groups, contains many examples of multicellularity – multicellularity by division in animals, multinucleate development in Fungi, and aggregative multicellularity in Dictyostelids, *Fonticula* and others (the distribution of multicellularity in Obazoa (Amorphea without Amoebozoa), is shown in Figure 1.2). This distribution of multicellularity was the basis of a theory that aggregative multicellularity may have been a precursor to multicellularity by division in Metazoans (Mikhailov et al. 2009, Dickinson et al. 2012). Under this theory, the ancestors of metazoans were aggregative multicellular organisms, which existed for a part of their life cycle as single cells. In the transition to multicellularity, the single cell section of the life
cycle was reduced to a single stage after gametes fuse, and instead of this large cell then dividing and releasing many zoospores (as in *Creolimax fragantissima*, an Ichthyosporan (Marshall et al. 2008)), the zoospores remained attached together, creating the first animal blastulas. Despite extensive cultivation, and in some cases genome sequencing, we do not have complete knowledge of the life cycles of many of the unicellular relatives of Metazoa. As well, superficial morphological and molecular similarities can be misleading. A key piece of evidence for the ancestral aggregative multicellularity theory
proposed by Dickinson et al. (2012) was the similarity in the arrangement of cells to make a polarized epithelium in dictyostelids and metazoans. While the structure and molecular components were indeed similar, careful phylogenetic analysis of the genes involved proved this to be a case of molecular convergence (Parfrey and Lahr 2013). Therefore there are multiple transitions to multicellularity within Amorphea.

There are several unicellular or colonial lineages of interest when studying the origin of multicellularity in animals (Fig. 1.2). Obazoa is a clade that includes the opisthokonts, and two unicellular groups – the apusomonads and the breviates (Brown et al. 2013). As Opisthokonta contains two examples of complex multicellularity – Fungi and Metazoa – the breviates and apusomonads could provide an important comparison for understanding these two very different types of multicellularity. Within Opisthokonta, Holozoa includes Metazoa and those unicellular organisms more closely related to them than to fungi (Fig. 1.2). Unicellular holozoans such as *Capsaspora* and *Sphaeroforma* have provided important insights into the origin of animal multicellularity (Sebé-Pedrós et al. 2013a, Fairclough et al. 2013, King et al. 2003, de Mendoza et al. 2013).

### 1.5 Aims of this Thesis

With the huge range of multicellular organization present in eukaryotes, the transitions from unicellular to multicellular lifestyles present inexhaustible scope for research. I present estimates of when various multicellular groups evolved using phylogenomics and relaxed molecular clocks in Chapter 2. In Chapter 3, I present the evolutionary history and features of unicellular versions of integrins, a cell adhesion molecule important for animal multicellularity.
Chapter 2: Timing the Origins of Multicellular Eukaryotes
Through Phylogenomics and Relaxed Molecular Clock Analyses

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Timing the origins of multicellular eukaryotes through phylogenomics and relaxed molecular clock analyses. In Ruiz-Trillo I, Nedelcu AM (eds) Evolutionary transitions to multicellular life: Principles and mechanisms. Springer Science+Business Media, doi: 10.1007/978-94-017-9642-2_1 ISBN 978-94-017-9641-5. Its contents have been amended here following the suggestions of the examination committee.
* these authors contributed equally to the manuscript

2.1 Abstract

Multicellularity has evolved many times during eukaryote evolution. Deciphering the evolutionary transitions to multicellularity requires a robust deep phylogeny of eukaryotes to clarify the relationships amongst multicellular groups and determine their closest unicellular relatives. Here I review progress in understanding of the phylogenetic relationships amongst multicellular and unicellular eukaryotes, as well as estimates of the ages of multicellular groups based on relaxed molecular clock (RMC) analyses. In addition, I present an RMC analysis of a large phylogenomic dataset to estimate the divergence dates of select major eukaryotic multicellular groups. My analyses (and other recent studies) tentatively suggest that multicellular eukaryotes such as Metazoa, Fungi and two of the major multicellular red algal taxa first emerged in the mid-Neoproterozoic, whereas the dictyostelids arose in the Paleozoic. I also hypothesize that the first multicellular organisms emerged within 300 – 600 Myr after the Last Eukaryotic Common Ancestor (LECA). The age of land plants is less clear and is highly dependent on methodology, the genes analyzed, and the nature of fossil constraints. In general, there is great variability in all these age estimates, and their credible intervals span hundreds of millions of years. These estimates are highly sensitive to both the models and methods of RMC analysis, as well as the manner in which fossil calibrations are treated in these analyses. As paleontological investigations continue to fill out the Proterozoic fossil record, genomic data is gathered from a greater diversity of eukaryotes and RMC
methodology improves, estimates of the ages of multicellular eukaryotes may converge to precise dates that can be correlated with Earth’s ancient geochemical record.

2.2 Introduction

2.2.1 Eukaryote Phylogeny

As knowledge of the deepest relationships between all extant eukaryotes improves, there is a better understanding of how multicellularity has developed several times in distantly related eukaryote lineages. Early phylogenies of eukaryotes based on small subunit ribosomal RNA genes (SSU rDNA) showed the Metazoa, plants, Fungi and many protistan groups emerging from an unresolved radiation. This radiation was preceded by the divergence of a series of protistan lineages, with anaerobic ‘amitochondriate’ protists emerging as the earliest branches next to the prokaryotic outgroup (Cavalier-Smith and Chao 1996). This supported the idea that eukaryotic evolution proceeded by a gradual increase in complexity, from simple cells without mitochondria, through more complex unicellular organisms, to complicated multicellular organisms. This understanding of eukaryotic evolution appeared to be founded on methodological artifacts (Roger 1999, Roger and Hug 2006) and on an incorrect notion of the nature of so-called ‘amitochondriate’ eukaryotic lineages. It is now clear that the latter all possess homologs of mitochondria in the form of mitochondrion-related organelles (MRO) (Tsaousis et al. 2012). Furthermore, because their SSU rDNA sequences have evolved more rapidly than those of other eukaryotes, their deep-branching position in the eukaryote tree is likely a result of the infamous long-branch attraction (LBA) artifact whereby they are artificially clustering with the long branches leading to the prokaryotic outgroup (Roger and Hug 2006). With more data from multiple genes and better analytical methods it has become clear that the apparently ‘deep-branching’ lineages on early SSU rDNA trees in fact emerge in multiple distinct places in the eukaryote tree (Roger 1999, Roger and Hug 2006, Keeling et al. 2005, Baldauf et al. 2000) as do multicellular groups that each show affinities to distinct ancestral unicellular protistan lineages (Burki et al. 2012).

Recent advances in our understanding of deep eukaryotic phylogeny have come from analyses of large sets of concatenated genes that provide more information on ancient nodes (Parfrey et al. 2010, Brown et al. 2012a, Burki et al. 2007). While the deep
branching order of eukaryotic lineages is still controversial (Zhao et al. 2012), a number of relatively ‘stable’ eukaryotic supergroups have been identified. One of the earliest recognized higher-level groupings of eukaryotes are animals, Fungi and their unicellular relatives (Wainright et al. 1993) collectively known as the Opisthokonta. In unrooted phylogenies of eukaryotes, opisthokonts are adjacent to the Amoebozoa, a group that includes a wide variety of unicellular amoebae, anaerobic species previously thought to be basal eukaryotes, and the Eumycetozoa (social amoebae, myxogastrids and relatives). Opisthokonts, Amoebozoa, in addition to Breviata, Apusomonadida and a number of unicellular organisms of unclear phylogenetic affiliation (Kim et al. 2006, Brown et al. 2013, Cavalier-Smith and Chao 2010) form the major division Amorphea (Adl et al. 2012), roughly equivalent to “unikonts” of Stechmann and Cavalier-Smith (2003) and Richards and Cavalier-Smith (2005). Excavata is a possibly paraphyletic supergroup including many long branches that contains several of the organisms originally thought to be basal to other eukaryotes, but are now known to be united by ultrastructural characteristics and molecular data (Simpson 2003, Simpson et al. 2006, Hampl et al. 2009). The remainder of eukaryotic diversity is encompassed in a grouping referred to as Diaphoretickes that contains most of the photosynthetic lineages of eukaryotes (Adl et al. 2012). Within Diaphoretickes, the Archaeplastida encompasses eukaryotes with a primary plastid, including glaucophytes, rhodophytes (red algae) and green algae (which includes land plants). SAR is an assemblage made up of the Stramenopiles, Alveolata and Rhizaria. These supergroups are generally well supported in phylogenomic analysis (Brown et al. 2013, Hampl et al. 2009, Burki et al. 2012), including the 159-gene analysis shown in Figure 2.1. Finally other lineages have more uncertain placements in the eukaryote tree including the haptophytes, cryptophytes, telonemids, and collodictyonids (Zhao et al. 2012, Burki et al. 2012).

It is uncertain how Diaphoretickes, Amorphea and Excavata are related, as this depends on the location of the root of the tree of all eukaryotes. Since the position recovered in early rDNA analyses has been discredited as an artifact of LBA, several other possibilities for the root have emerged, and rare genomic changes have been used to define its location. For example, a fusion between dihydrofolate reductase (DHFR) and thymidylate synthase (TS) (Stechmann and Cavalier-Smith 2002), and the distribution of
myosin II, suggested that the root might lie between Opisthokonts+Amoebozoa (Amorphea) and all other eukaryotes (Richards and Cavalier-Smith 2005). More recently, it has become clear that the distribution of these features in various eukaryote lineages could not be simply explained by ‘single gain’ scenarios and that so-called ‘rare’ changes may have occurred more frequently than was once thought (Kim et al. 2006, Roger and Simpson 2009). Using a molecular phylogenetic approach, Derelle and Lang (2012) analysed a collection of mitochondrion-derived genes and found support for a root between Amorphea other eukaryotes. However, analyses of another dataset with better taxonomic sampling (He et al. 2014) supported a eukaryotic root between Discoba (a group within Excavata) and all other eukaryotes (lack of data for other Excavates meant this study could not conclude whether Excavates was a monophyletic group). Other root positions have been recovered using other types of data. For example, an approach minimizing gene family duplication and loss apparently supported a root between Opisthokonts and other eukaryotes (Katz et al. 2012). In contrast, Cavalier-Smith has suggested that the root lies between the Euglenozoa and all other eukaryotes because the former lack a number of molecular and morphological features that are conserved in most eukaryotes (Cavalier-Smith 2012). Clearly, there is no consensus on the position of the eukaryote root and many candidate positions are plausible given the current evidence.

2.2.2 Placing Multicellular Groups on the Tree of Eukaryotes

Multicellularity represents a spectrum of organization ranging from simple colonies of cells to complex differentiated multicellular organisms (Bonner 1998). Colonial growth is common throughout eukaryotes and varies in complexity and the degree of intercellular integration. Examples include diatoms, which can interlock their silica shells to form chains, and the ciliates of the genus Zoothamnium, which have a sessile colonial form that can contract when exposed to stimuli (Bonner 1998). In this chapter, I do not consider simple colonial organisms; instead I focus on more complex multicellular taxa (e.g., Metazoa, Fungi and land plants (Embryophyta), as well as less well-known forms such as red algae and dictyostelids). Many of these possess sophisticated mechanisms for the communication and transport of nutrients between cells. It is generally accepted that most of these multicellular groups evolved independently from distinct unicellular ancestors (Knoll 2011, Brown and Silberman 2013).
The largest and/or most familiar multicellular organisms develop through cells dividing but not separating, followed by the differentiation of cell lineages into different specialized types. This kind of ‘multicellularity-by-division’ is employed in both simple organisms with a few cell types (e.g., the green algal genus *Volvox*) and in complex organisms with hundreds of cell types (e.g., Metazoa) (Rokas 2008). Alternatively, multicellular organisms known informally as ‘cellular slime moulds’ can develop through the aggregation of single cells (Brown and Silberman 2013). In this form of simple multicellularity, organisms exist as single cells for part of their life cycle, but come together to form specialized organs for the distribution of spores. While the same basic challenges of cell adhesion and communication need to be solved for both types of multicellularity, they appear to have evolved in different types of environments. Aggregative multicellularity has evolved in organisms that live predominantly in terrestrial environments, while multicellularity-by-division has apparently evolved in lineages which were originally aquatic (Bonner 1998).

Multicellular organisms face several challenges that are not relevant to strictly unicellular organisms, including cell-to-cell communication and adhesion. The various distinct lineages of multicellular organisms have solved these challenges in different ways. To make inferences about the transition to multicellularity both how these multicellular groups are related to each other and to unicellular organisms, and how the genes involved in their multicellularity have evolved must be understood. For example, Dickinson and colleagues have argued that biochemical and morphological similarities between the epithelial tissues of one group of aggregative multicellular organisms (dictyostelids) and the Metazoa indicated their common amorphean ancestor was multicellular (Dickinson et al. 2012). However, Parfrey and Lahr (2013) conducted more detailed evolutionary bioinformatic analyses that show the proteins involved are paralogs that within a larger protein family, and this family is commonly found in unicellular amorpheans. The molecular similarities are a result of convergent evolution in Dictyostelids and Metazoa when these proteins were co-opted for functions related to multicellularity. Dictyostelid and metazoan epithelia are therefore unlikely to be homologous (Parfrey and Lahr 2013). In general, there is currently little reason to
suppose that any of the molecular mechanisms underpinning multicellularity in the various lineages I discuss are homologous.

Multicellularity-by-division is exhibited by multiple lineages within the tree of eukaryotes. Within Archaeplastida (Figure 2.1) this form of multicellularity has evolved both within the red algae and green algae. Both Florideophyceae and Bangiales within the red algae are multicellular, and a fossil Bangia-like organism, *Bangiomorpha*, apparently provides the earliest fossil evidence of eukaryotic multicellularity (i.e., dated at 1198 +/- 24 Ma) (Butterfield 2000). Within the green algae, multicellularity has developed multiple times, giving rise to a wide variety of forms, such as the nets of *Hydrodictyon* (Bonner 1998) and the ball-like volvocine algae (Herron 2009). *Volvox* is a well-studied example of the latter, which is useful in a comparative genomics context because many close relatives with varying degrees of complexity exist (Herron 2009, Kirk 2005). Finally, the Embryophyta (land plants) are the best-known and most conspicuous multicellular organisms within the Archaeplastida.

Animals (Metazoa) include the most complex of all multicellular organisms, and range from simple sponges to mammals with elaborate nervous systems. Besides animals, the supergroup Opisthokonta includes another well-known multicellular-by-division group, the Fungi. Fungi often display filamentous ‘hyphal’ growth and, complex multicellularity with tissue differentiation occurs in the fruiting bodies of several lineages that evolved separately in the Basidiomycota and Ascomycota (Knoll 2011, Stajich et al. 2009). Other fungal organisms have apparently experienced reductive evolution to unicellularity (e.g., yeasts) (Stajich et al. 2009). Distantly related eukaryotes, the Oomycetes (a stramenopile lineage) have converged on a similar lifestyle to that of Fungi and show filamentous growth (Beakes et al. 2012). Finally, brown algae, another stramenopile group, include large kelps and the model organism *Ectocarpus* (Cock et al. 2010).
Among organisms showing the second main type of multicellularity (i.e., aggregative multicellularity), the best studied are the dictyostelids. Many other organisms that display aggregation have been discovered over the years, but only recently have they been placed in the tree of eukaryotes using molecular data (Brown and Silberman 2013). The dictyostelids belong to the supergroup Amoebozoa, and are closely related to the plasmodial slime molds, which achieve a macroscopic form by growing into multinucleate plasmodia (Schilde and Schaap 2013). Other aggregative protists include *Copromyxa*, another amoebozoan (Brown et al. 2011), *Capsaspora owczarzaki*, a relative of Metazoa (Sebé-Pedrós et al. 2013b), and *Fonticula alba*, an amoeba that groups as a sister lineage to Fungi (Brown et al. 2009). Within the SAR clade, there are examples of aggregative multicellularity within each of the three main groups: *Guttulinopsis* within Rhizaria (Brown et al. 2012a), *Sorodiplophrys* in Stramenopiles (Dykstra and Olive 1975), and the ciliate *Sorogena* within Alveolates (Olive and Blanton 1980). Finally, the acrasid amoebae within excavates are aggregative (Brown et al. 2012b), leaving Archaeplastida as the only supergroup without an aggregative multicellular representative.

### 2.2.3 Using Molecular Data to Date the Emergence of Lineages

As well as being invaluable for the elucidation of phylogeny, molecular data can be used to estimate the date of divergence between organisms. From molecular sequence data it is possible to calculate an evolutionary distance, which is the product of rate of substitution (i.e., fixed nucleotide or amino acid changes) and time. Consequently, if the date of divergence for two taxa is known, an average rate of substitution can be inferred. If the rates of substitution are equal across all branches on a tree (i.e., a ‘molecular clock’ holds), this rate can then be used to convert branch lengths from the rest of the tree to dates (Zuckerkandl and Pauling 1965). Given that evolutionary rates often vary across subgroups of the tree of life (for example, see discussion of LBA above), methods using strict molecular clocks are not appropriate unless explicit tests are conducted to prove that their use is justified (Welch and Bromham 2005, Takezaki et al. 1995).

Once it became clear that a strict molecular clock does not generally hold (Langley and Fitch 1974), many efforts were made to ‘relax’ the molecular clock (Sanderson 1997, Sanderson 2002, Kishino et al. 2001, Thorne et al. 1998, Lepage et al. 2007). Strategies ranged from assigning subsections of the tree to evolve at different rates (Yoder and...
Yang 2000), to employing a complex Bayesian framework to model branch-specific rates, along with other parameters associated with the tree (Yang 2006). Substitution rates depend on biological processes such as mutation rate and generation time (Ho 2009), and so may be correlated on neighbouring branches. Whether the correlation is applicable between branches that cover a large phylogenetic distance such as all of eukaryotes is less clear. Correlated models (such as the lognormal (LogN) (Kishino et al. 2001) and Cox-Ingersoll-Ross (CIR)(Lepage et al. 2007) used in my analyses below) draw the rate on a particular branch from a probability distribution of rates centered on the rate of the ‘parent’ branch. Uncorrelated models (for example, the uncorrelated gamma (UGam) model in my analyses) draw rates for each branch from one global probability distribution, not taking into consideration the rates on adjacent branches (Drummond et al. 2006).

Another difficulty with molecular clock analysis concerns the assignment of fossil-based time calibrations (Benton and Donoghue 2007, Parham et al. 2012). There is only indirect evidence for the date of divergence of two species: groups under study must have existed for some time before the right conditions occurred for preservation in the geological record. Moreover, even if an organism fossilizes, estimates of its geological age also have associated uncertainty. In addition, identification of fossils is often controversial (especially in the case of simple ‘soft-bodied’ multicellular or unicellular organisms), as they often lack characteristic features that would allow them to be definitively assigned to an extant group. Furthermore, fossil assignments depend on whether a fossil is a member of the ‘crown group’ (i.e., it descends from the most recent common ancestor (MRCA) of all extant species in the group), or instead represent a ‘stem group’ lineage, (i.e., it diverged prior to the MRCA of extant members of the group, and thus does not possess any currently living descendant) (Benton and Donoghue 2007). Unfortunately, it is sometimes difficult to determine whether a given fossil corresponds to an extinct crown group lineage or is a stem group organism. Clearly, this distinction is extremely important as each of these types of fossils conveys different information about the ages of nodes on molecular phylogenies of extant organisms. For example, stem lineage fossils may be older or younger than the node defining the MRCA of the crown
group on phylogenies, whereas crown group fossils must be younger than this node (and can therefore provide a lower bound on its age).

In addition to the difficulties associated with assigning fossil constraints to specific nodes on phylogenetic trees, another problem relates to the fact that crown group fossil dates must always be treated as minimum possible ages (as the MRCA of the group must predate the appearance of the first fossil). While a few groups of protists have a continuous fossil record that can be fairly reliably translated into minimum and maximum dates (Berney and Pawlowski 2006), most organisms on the tree of life are only sporadically fossilized, and so a fossil represents only a minimum restriction on the age of that group, which is not particularly informative for subsequent analyses.

Finally, there has been much debate over how fossil dates should be treated during RMC analyses, notably because of the uncertainty associated with the dating of the rocks in which the fossils are found (for a detailed review on this question, see (Parham et al. 2012) and references therein). There are several ways of applying fossil constraints (Ho and Phillips 2009). Treating calibrations as ‘hard bounds’ implies that the estimated age of a constrained node will necessarily fall within the fixed time intervals specified by the paleontological evidence (Kishino et al. 2001). In contrast, the ‘soft bound’ approach uses a probabilistic treatment of fossil age data, and therefore can accommodate potential error in calibrations. The simplest of these is the use of a uniform prior probability distribution for the node falling within the bounds defined by the fossil age data (Yang and Rannala 2006), leaving some probability of the node falling outside the bounds (e.g., a smoothly decreasing probability distribution on each side of the bounds, as implemented in Phylobayes (Lartillot et al. 2009) and used for my analyses below). Alternatively, fossil evidence can also be represented as parametric probability distributions such as lognormal or gamma, with a ‘mix’ of a hard lower bound and soft upper bound (e.g., as implemented in the software BEAST (Drummond et al. 2006)). The manner in which fossil dates are treated by the software that is estimating dates can have a large impact on the final results (Inoue et al. 2010).

Translating fossil and other geological evidence to a range or distribution of dates is often subjective and has attracted much criticism (Shaul and Graur 2002). Recently, efforts have focused on formalizing strategies for interpreting fossil data (Parham et al.
and selecting age distributions (Nowak 2013); however, these have yet to be widely applied.

2.3 Methods

2.3.1 Dataset Construction

In collaboration with Dr. Matthew Brown, I updated a previously developed data set of 159 aligned proteins from diverse eukaryote lineages (Brown et al. 2013) to provide the most possible calibrated nodes based on available fossil evidence (total of 85 taxa). BLAST was used to search reference genes against all protein data available for a specific taxon. The top hit in each taxon was then searched using BLAST against the ORTHOMCL database of orthologous protein families (www.orthomcl.org), and discarded if it did not match the correct orthologous group for the gene of interest. Single gene trees were constructed, and an in house script was used to identify taxa branching with high support that contradict well-established phylogenetic groupings. When manual examination of the single-gene trees revealed possible lateral gene transfer or contamination (e.g. a single taxon from one established group branching with high support with several taxa from a different established group) the sequence for that gene and taxa were removed. Each gene was then re-aligned, trimmed to remove ambiguously aligned sites using BMGE (Criscuolo and Gribaldo 2010), and concatenated to form the final 43099 site alignment.

2.3.2 Reference Phylogeny

A maximum likelihood (ML) phylogenetic tree was obtained from 60 heuristic searches employing RAxML version 7.2.6 (Stamatakis 2006) under the LG + Γ +F amino acid substitution model. Bootstrap support (BS) for splits was estimated from 500 pseudoreplicates. To assess whether the reference phylogeny was robust to methodological bias, a Bayesian estimation of the tree was also performed. Four chains of Markov Chain Monte Carlo (MCMC) were run with Phylobayes 3.2 (Lartillot et al. 2009) for each of the CAT-GTR, CAT-POISSON and catfix C60-Poisson substitution models. For each of the three substitution models, the four MCMC chains did not
converge, although the post-burnin consensus tree was identical to the ML tree except for an unresolved multifurcation at the base of Excavata.

### 2.3.3 Fossil Calibrations

Fossil calibrations (Table 2.1) were taken from (Parfrey et al. 2011) with adjustments described below. Four calibrations (Gonyaulacales, spirotrichs, Foraminifera and euglenids) were removed due to insufficient sampling of taxa from the clade of interest. The ‘ciliate’ tetrahymenol-based calibration (Summons and Walter 1990) was removed, as the biomarker used, tetrahymenol, is also found in aneorobic protists (Takishita et al. 2012). Seven calibrations were adjusted by using a different taxa of equivalent phylogenetic position for which genomic data was available. As insufficient data was available from *Isochrysis galbana* or any equivalent taxa within the haptophytes, the coccolithophore calibration was adjusted from a narrow limit, to a wide range with an uninformative maximum (3000 Ma). A recent discovery of the oldest known cestode (tapeworm) (Dentzien-Dias et al. 2013) was used to add a calibration for Platyhelminths. The minimum age (250 Ma) was taken from the youngest possible age of the fossil (lower boundary of the Permian) and the upper boundary was taken from the limit on the next-oldest calibrated node (Bilateria). With the ten remaining calibrations unchanged, nineteen calibrations were used to date the phylogeny.

### 2.3.4 Relaxed Molecular Clock Analyses

The ML phylogeny was used for relaxed clock molecular dating analyses with Phylobayes 3.2 (Lartillot et al. 2009). For all analyses, a birth-death tree prior was applied, and a discrete gamma distribution was used to model among site rate variation. Two chains were run until diagnostic statistics from the ‘tracecomp’ function of Phylobayes indicated convergence or until estimated dates on nodes of interest for the two chains were < 10% different.

As it is unclear which models of evolution are most realistic, I present results obtained with three different RMC models, two different substitution matrices, and two ways of treating fossil calibrations, for a total of twelve sets of parameters. The RMC models included both uncorrelated (UGam) and correlated models (LogN, CIR), where

**Table 2.1 Calibrations used for relaxed molecular clock analysis. (following page)**
<table>
<thead>
<tr>
<th>Taxon Fossil</th>
<th>Node Specification</th>
<th>Min (Ma)</th>
<th>Max (Ma)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pennate diatoms</strong></td>
<td><em>Phaeodactylum</em> / <em>Thalassiosira</em></td>
<td>80</td>
<td>110</td>
<td>(Kooistra et al. 2007)</td>
</tr>
<tr>
<td>Oldest pennate</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Eudicots</strong></td>
<td><em>Arabidopsis</em> / <em>Oryza</em></td>
<td>125</td>
<td>133.9</td>
<td>(Friis et al. 2010)</td>
</tr>
<tr>
<td>Eudicot pollen</td>
<td></td>
<td></td>
<td></td>
<td>(Sun et al. 2011)</td>
</tr>
<tr>
<td><strong>Angiosperms</strong></td>
<td><em>Arabidopsis</em> / <em>Welwitschia</em></td>
<td>133.9</td>
<td>425</td>
<td>(Crane et al. 1994)</td>
</tr>
<tr>
<td>Oldest angio pollen</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td><strong>Diatoms</strong></td>
<td><em>Pelagophyceae</em> / <em>Thalassiosira</em></td>
<td>133.9</td>
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<td>(Harwood et al. 2007)</td>
</tr>
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<td></td>
<td></td>
</tr>
<tr>
<td><strong>Coccolithophores</strong></td>
<td><em>Emiliania</em> / <em>Prymnesium</em></td>
<td>203.6</td>
<td>3000</td>
<td>(Bown 1998)</td>
</tr>
<tr>
<td>Earliest</td>
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<tr>
<td>Heterococcolith</td>
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<tr>
<td><strong>Dinoflagellates</strong></td>
<td><em>Alexandrium</em> / <em>Symbiodinium</em></td>
<td>240</td>
<td>300</td>
<td>(Fensome et al. 1999)</td>
</tr>
<tr>
<td>Earliest gonyaulacales</td>
<td></td>
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<tr>
<td><strong>Platyhelminths</strong></td>
<td><em>Taenia</em> / <em>Schistosoma</em></td>
<td>250</td>
<td>555</td>
<td>(Dentzien-Dias et al. 2013)</td>
</tr>
<tr>
<td>Earliest cestode</td>
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<tr>
<td><strong>Endopterygota</strong></td>
<td><em>Apis</em> / <em>Drosophila</em></td>
<td>284.4</td>
<td>350</td>
<td>(Dostál and Prokop 2009)</td>
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<tr>
<td>Mecoptera</td>
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<tr>
<td><strong>Amniota</strong></td>
<td><em>Gallus</em> / <em>Homo</em></td>
<td>328.3</td>
<td>400</td>
<td>(Smithson and Rolfe 1990)</td>
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<tr>
<td>Westlonthania</td>
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<tr>
<td><strong>Ascomycetes</strong></td>
<td><em>Schizosaccharomyces</em> / <em>Coprinus</em></td>
<td>400</td>
<td>1000</td>
<td>(Taylor et al. 1999)</td>
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<tr>
<td>Paleopyrenomycites</td>
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<tr>
<td><strong>Trachaeophytes</strong></td>
<td><em>Physcomitrella</em> / <em>Arabidopsis</em></td>
<td>425</td>
<td>471</td>
<td>(Kenrick and Crane 1997)</td>
</tr>
<tr>
<td>Earliest trachaeophytes</td>
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<tr>
<td><strong>Embryophytes</strong></td>
<td><em>Mesostigma</em> / <em>Oryza</em></td>
<td>471</td>
<td>600</td>
<td>(Rubinstein et al. 2010)</td>
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<tr>
<td>Land plant spores</td>
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<tr>
<td><strong>Vertebrates</strong></td>
<td><em>Branchiostoma</em> / <em>Homo</em></td>
<td>520</td>
<td>555</td>
<td>(Shu et al. 1999)</td>
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<tr>
<td>Haikouichthys</td>
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<tr>
<td><strong>Bilateria</strong></td>
<td><em>Branchiostoma</em> / <em>Caenorhabditis</em></td>
<td>555</td>
<td>630</td>
<td>(Martin 2000)</td>
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<tr>
<td>Kimberella</td>
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<tr>
<td><strong>Arcellinida</strong></td>
<td><em>Acanthamoeba</em> / <em>Copromyce</em></td>
<td>736</td>
<td>3000</td>
<td>(Porter et al. 2003)</td>
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<tr>
<td>Paleoarcella</td>
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<tr>
<td><strong>Animals</strong></td>
<td><em>Amphimedon</em> / <em>Homo</em></td>
<td>632</td>
<td>3000</td>
<td>(Love et al. 2009)</td>
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<tr>
<td>LOEMs, sponge biomarkers</td>
<td></td>
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<td>(Cohen et al. 2009)</td>
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<tr>
<td><strong>Chlorophytes</strong></td>
<td><em>Volvox</em> / <em>Acetabularia</em></td>
<td>700</td>
<td>3000</td>
<td>(Butterfield et al. 1994)</td>
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<td>Palaeastrum</td>
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<tr>
<td><strong>Florideophyceae</strong></td>
<td><em>Chondrus</em> / <em>Porphyra</em></td>
<td>550</td>
<td>3000</td>
<td>(Xiao et al. 2004)</td>
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<td>Doushantuo red algae</td>
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<tr>
<td><strong>Red algae</strong></td>
<td><em>Porphyra</em> / <em>Cyanidioschyzon</em></td>
<td>1174</td>
<td>3000</td>
<td>(Butterfield 2000)</td>
</tr>
<tr>
<td>Bangiomeropa</td>
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</table>
correlated models indicate that the rate of one branch is dependent on the rate of the parent branch. The uncorrelated gamma multipliers RMC model (UGam, Drummond et al. (2006)) takes rates from a single gamma distribution, which is fitted to the data during Bayesian inference. The correlated lognormal RMC model (LogN) (Thorne et al. 1998) allows variation of rates following a lognormal distribution. The Cox-Ingersoll-Ross (CIR) model (Lepage et al. 2007) is correlated from branch to branch as well. Under this model changes near the mean rate are considered likely, but larger or additive changes are penalized, pushing the value back to the mean. Two substitution matrices were considered. The site-heterogeneous empirical profile mixture model C60-Poisson (C60) (Le et al. 2008) considers all amino acid substitutions as equally likely, but possible amino acids at a site are defined by sixty empirically derived site classes. The LG substitution matrix (Le and Gascuel 2008) uses an empirically defined matrix of probabilities of transition from one amino acid to another, and does not have different site classes. Two ways of treating fossil calibrations were used - “soft” and “hard” bounds. Under the hard bound parameter, the age of that node cannot fall outside of the range defined by the calibration. Soft bounds allow a 5% probability that the node is either older or younger than the specified range.

Dating evolutionary events requires the specification of the position of the root of the eukaryotic tree. For this reason, three alternative positions for the root were tested: (i) between Amorphea and the remainder of eukaryotes (i.e., the classical “unikont-bikont” rooting (Derelle and Lang 2012); (ii) between the Obazoa (Opisthokonta + apusomonads + breviates) and all other taxa (Katz et al. 2012); and (iii) between Excavata and all other eukaryotes (He et al. 2014). Interestingly, these three alternative placements had little impact on the estimated ages of the various groups considered here (discussed further in Eme et al. (2014)). Therefore I only discuss our RMC age estimates for major multicellular groups obtained with the Amorphea root.

2.4 Results and Discussion

In the last few decades, many attempts have been made to date divergences deep within the tree of life, including many of the multicellular eukaryote lineages. Here I review the most recent attempts that employ relaxed molecular clock methods. In
addition I present an analysis of the age of various multicellular groups based on a phylogenomic dataset with molecular data and fossil calibrations from representatives of the full breadth of eukaryotic diversity. I focus specifically on multicellular groups that are well represented in my own analyses (Figure 2.1 and below) including the Metazoa, Fungi, embryophytes (land plants), the red algae and dictyostelids.

For my estimation of the ages of multicellular groups (nodes indicated by letters in Figure 2.1), I used the Bayesian implementation of RMC models in Phylobayes (Lartillot et al. 2009) to analyze a large phylogenomic dataset (159 proteins, 85 taxa). The phylogenetic tree was obtained using maximum likelihood analysis and was calibrated by nineteen fossil calibrations from diverse eukaryote groups, the majority of them taken from Parfrey et al. (2011). My analyses included three different RMC models (the uncorrelated UGam model, and the correlated LogN and CIR models), two substitution matrices (the site-heterogeneous empirical profile mixture model C60-Poisson (C60)(Le et al. 2008) and the more classical site-homogeneous LG substitution matrix (Le and Gascuel 2008)), and two ways of treating fossil calibrations ("soft" and "hard" bounds).

Figure 2.2 Estimates of the age of the most recent common ancestor of Metazoa (a), Fungi (b) and Dictyostelids (c). Results are shown for the combination of three different RMC models (UGam, LogN and CIR) with two substitution models (C60, squares; LG, circles), and with soft (filled shapes) or hard (open shapes) bounds. Error bars indicate 95% credible intervals. For each node with calibration, the error bar above ‘Cal’ shows the age range used to calibrate the node. Where upper bound is not shown, it was equal to 3000 Ma. Asterisks (*) indicate estimates where the two chains showed more than a 10% difference in either the mean, the upper 95% limit or the lower 95% limit.
A few general trends can be noted in my results. First, soft bounds yielded younger date estimates than hard bounds when other parameters (i.e., relaxed clock model, substitution matrix) were identical (Figures 2.2 and 2.3). As the use of hard bounds is hardly justified given the uncertainties discussed above, I will mainly focus on results obtained with soft bounds. Second, the uncorrelated UGam relaxed molecular clock model gave larger credible intervals (so that the range for soft and hard bounds largely overlapped), with little change based on the substitution model. Third, analyses with the LogN and CIR rate evolution models showed more variation, both in terms of estimated ages of nodes, and of size of the confidence interval. This variation was often only seen when both the type of calibration used (hard or soft bounds) and the substitution matrix (C60 or LG) was changed, underlying the complex interaction between the various features of RMC analyses in influencing the age estimates obtained. Fourth, in general, the choice of the substitution matrix often did not make much of a difference (Figure 2.2 and 2.3, compare black squares to black circles for a given clock model), although more substitution matrix-based changes were seen in CIR analyses. For further discussion on the impact of these various parameters, see Eme et al. (2014).

2.4.1 Metazoa

There have been a large number of molecular clock-based studies attempting to date groups within Metazoa, in particular the Bilateria. These have often sought to illuminate the Cambrian explosion, an apparently sudden emergence of many divergent bilaterian phyla in the fossil record at the start of the Cambrian Period (530 Ma)(Levinton 2008). While much has been made about this change in the fossil record, molecular clock analyses often date the first Bilaterian divergence as much older (Hedges et al. 2004, Erwin et al. 2011). Basal metazoans such as cnidarians, placozoans and sponges existed for many years previous to this, as sponge biomarkers (Love et al. 2009) and fossil evidence for sponges (Cohen et al. 2009) dates to 632 Ma. The transition from unicellularity to multicellularity would have happened before the last common ancestor of all extant Metazoans (Figure 2.1, node a) and I focus on the estimated age of this node in the previous studies.

Hedges et al. (2004) estimated the Porifera-Animalia divergence at 1,351 ± 120 Ma (1116–1586), more than twice as old as the first fossil evidence for Metazoa. Their 17 –
19 protein gene dataset (depending on the method applied) was calibrated using the bird-mammal divergence as a primary calibration, and three additional secondary calibrations; various molecular clock methods were used, including Bayesian RMC modeling. Secondary calibrations involve using a date obtained from one molecular clock analyses as a calibration for another molecular clock calculation, a practice which has been vigourously criticized (Graur and Martin 2004) for failing to add new information to the dating analyses, propagating earlier biases from the analyses that generated the calibrations, and for failing to properly ‘carry forward’ the uncertainty associated with them.

A much younger estimate was obtained with analyses using nine primary invertebrate fossil calibrations and a dataset of seven genes (Peterson and Butterfield 2005). These molecular clock analyses (Sanderson 2003) yielded different estimates depending on the phylogenetic method used to construct the tree. The minimum evolution tree yielded an estimate of 664 Ma, whereas the ML tree gave 867 Ma. Later reanalyses of this dataset (Roger and Hug 2006) found substantial variation in the estimates (approximately 750 – 1300 Ma) and extremely broad confidence intervals depending on the substitution models, relaxed molecular clock method/models and the manner in which fossil constraints were applied.

Using 22 Phanerozoic minimum calibrations with only four maximum calibrations based on fossils from a wide diversity of eukaryotic groups, Berney and Pawlowski (2006) performed Bayesian RMC analyses of SSU rDNA sequences to date the divergence of major eukaryotic groups. They estimated that the primary split within Metazoa occurred 812 Ma (671 – 985).

In contrast to Berney and Pawlowski’s analyses of one gene with many fossil calibrations, Lartillot et al. (2009) analyzed a many-gene dataset (68 genes) from 52 Holozoons (Metazoa plus unicellular opisthokont relatives) using only two fossil calibrations (within the Metazoa) as soft bounds. Dates were estimated using the PhyloBayes 3 software, with a CAT-GTR substitution model and lognormal autocorrelated RMC model. An age of ~800 Ma (700 – 900) was obtained for the MRCA of Metazoa.
With the goal of estimating the age of eukaryotes and their supergroups, Parfrey et al. (2011) analyzed a 15-gene dataset with rich taxon sampling and 22 fossil calibrations associated with different nodes throughout eukaryotic diversity. Using the uncorrelated lognormal RMC model in BEAST (Drummond et al. 2006), they found the age of the last common ancestor of all Metazoans to be 780 Ma (782 – 820). The estimate was about 60 Myr younger when the seven Proterozoic fossil calibrations (including Bangiomorpha) were excluded.

In a taxon-rich analysis focused on Metazoan evolution, Erwin et al. (2011) analyzed a 7-gene dataset with 118 metazoan taxa and 8 other taxa from Amorphea, with 24 fossil calibrations. Their Phylobayes analyses estimated the common ancestral Metazoan node to be between 747 (690 – 825) and 1093 (962 – 1260) Ma, depending on tree topology, RMC model (CIR or UGam), prior probability on the age of the root and the degree of soft bound relaxation (i.e., ranging from 5-50% prior probability of the node falling outside of the interval defined by the fossil dates).

While the geological evidence supports a bound on the youngest possible age for animals at 632 Ma, there is lack of clear evidence for a plausible upper bound. Therefore, for my analyses, I employed an uninformative maximum age for this node at 3000 Ma, following Parfrey and colleagues (Parfrey et al. 2011). All of my analyses placed the last common ancestor of Metazoa within 220 Myr of the lower calibration, with age estimates between 698 Ma and 851 Ma, depending on the parameters used (Figure 2.2a). These estimates fall within the range of age estimates of most of the other recent studies discussed above; these span roughly 650 – 850 Ma (Peterson and Butterfield 2005, Berney and Pawlowski 2006, Parfrey et al. 2011). The exceptions are the estimates obtained by Erwin and colleagues (2011). They recovered slightly older estimates (i.e., ~750 – 1100 Ma) that appeared to be fairly strongly influenced by the root age prior placed on the Holozoae-Fungi split.

2.4.2 Fungi

Complex multicellularity with tissue differentiation in fruiting bodies has evolved independently within two subgroups of the Fungi: the ascomycete lineage Pezizomycotina and the basidiomycete lineage Agaricomycotina (Stajich et al. 2009). Here I will focus specifically on the origin of simple filamentous ‘hyphal’ growth, which
was likely a feature of the common ancestor of most extant Fungi (with the possible exception of Cryptomycota and Microsporidia) (Stajich et al. 2009). Although the deepest branching order of the fungal tree is still controversial, I will provisionally consider the Chytridiomycota and Blastocladiomycota to form a clade that split basally from all other Fungi (Ebersberger et al. 2012, James et al. 2013), excluding the Cryptomycota and Microsporidia.

There have been few efforts to apply molecular clock methods to the Fungi as a whole, or even to specific groups of fungi, largely because of the lack of fungal fossils, and the difficulty of classifying the fossils that do exist. Using the Langley-Fitch method from the r8s software, Taylor and Berbee (2006) showed that age of the node uniting \textit{Rhizopus} (Mucoromycotina) and the Ascomycota + Basidiomycota group was highly uncertain (estimates ranged from 435 – 1979 Ma depending on the calibration used) (Taylor and Berbee 2006). When three calibrations (one each within Fungi, Metazoa and plants) were used, an age of 792 Ma was estimated. More recently, they used Bayesian analyses implemented in the BEAST software program to analyze a 50-gene dataset with three primary calibrations and a prior on the age of opisthokonts (Berbee and Taylor 2010). These analyses led to large confidence intervals for the majority of nodes, including the \textit{Rhizopus}/Ascomycota+Basidiomycota split which they dated at ~ 750 Ma. Global eukaryotic analyses dated the deepest split in extant Fungi (i.e., basal Chytridiomycota versus all other Fungi, a deeper divergence than the one considered by Berbee and Taylor (2010)) at 798 Ma (634 – 1003) (Berney and Pawlowski 2006) and 1070 Ma (980 – 1220) (Parfrey et al. 2011).

In my analyses, the CIR model showed the greatest range of age estimates for the split of the Chytridiomycota+Blastocladiomycota group from all other fungi (Figure 2.1, node b). It yielded both the youngest age estimate (744 Ma, employing LG + soft bounds), and the oldest (1152 Ma, employing C60 + hard bounds, Figure 2.2b). This range illustrates the difficulty of separating the synergistic effects of different parameters in molecular clock analysis. CIR also recovered wide credible intervals on the age, whereas the LogN model had the smallest (and non-overlapping) intervals, with hard bound estimates significantly older than the soft bound estimates. Soft bound estimates for the age of Fungi were similar (~760 Ma) over all combinations of substitution and RMC.
models, with the exception of the CIR + C60 combination, which was notably older (920 Ma).

My estimates for the age of Fungi are consistent with those of Berney and Pawlowski (2006). Surprisingly, Parfrey et al. (2011) estimated an older age (980 – 1,220 Ma), despite the fact most of their calibrations were identical to those used in this study. The paucity of fossil constraints within Fungi (i.e., only one) is likely largely responsible for this discrepancy; because this node is not strongly constrained by ancestral or descendant nodes, it is more susceptible to age estimate variation depending on the dataset and methodologies used.

### 2.4.3 Dictyostelids

Using either a six-protein dataset or an rDNA tree, Fiz-Palacios et al. (2013) estimated the divergence of dictyostelids using a combination of four fossil calibrations from opisthokonts and land plants. Variation in estimates was found between those obtained by the Bayesian package BEAST (employing the uncorrelated lognormal RMC model) and MCMCtree (using independent rates RMC model), with most estimates centering between 570 – 730 Ma. The one lower estimate obtained by Fiz-Palacios and colleagues (341 Ma (247 – 699)) was obtained when only the two opisthokont fossils were used and among site rate variation (ASRV) was modeled using a gamma distribution (modeling ASRV is essential for model realism (Yang 1996, Roger and Hug 2006)). Their analyses showed that trees calibrated with only the two plant fossils failed to return appropriate dates for the opisthokont fossils using fossil cross-validation (Near and Sanderson 2004). Land plants are phylogenetically distant from Amoebozoa, and so in absence of better sampling of fossil calibrations across the tree, using only plant fossils may heavily distort the results.

My estimates for the age of the last common ancestor of dictyostelids were around 330 Ma (Figure 2.2c) and remained similar across most clock models and substitution matrices (with soft bounds). However, even when restricting attention to soft bounds analysis, confidence intervals tended to be very wide, spanning from 160 to 430 Myr. In contrast, age estimates obtained by Fiz-Palacios and colleagues (2013) were much greater (~530 – 730 Ma), except in the single case described above where their estimate was closer to ours (~340 Ma).
Figure 2.3 Estimates of the age of the most recent common ancestor of Stomatophyta and Streptophyta (a), and Bangiales/Florideophyceae and Rhodophyceae (b). See Figure 2.2 for details.
2.4.4 Embryophytes, Stomatophytes and Streptophytes

Embryophytes, the ‘land plants’, emerge from within the Streptophyta clade that also includes a number of organisms with unicellular, colonial/filamentous and multicellular forms (Leliaert et al. 2012). Multicellularity seems to have evolved multiple times within the streptophytes, and reversion to simpler unicellular forms has probably occurred as well (Becker and Marin 2009, Leliaert et al. 2012). It is therefore difficult to pinpoint the origins of multicellularity within this group, although a key aspect of ‘complex multicellularity’ associated with embryophytes is the development of a diploid zygote into a multicellular diploid sporophyte (Becker and Marin 2009). Here I will focus on the crown embryophyte node (uniting liverworts with all other land plants), which post-dates the origin of embryophyte-type multicellularity and the crown streptophyte node which predates it. Note that many recent molecular clock studies (including ours below) have not included liverworts (the deepest branching embryophytes), so the crown ‘land plant’ node referred to in these studies is the somewhat younger common ancestor of the Stomatophyta (Figure 2.1, node e) that includes mosses and all other land-plants (Clarke et al. 2011). Fossil constraints that are commonly included in molecular clock analyses correspond to the first appearance of crown stomatophyte and embryophyte spores at ~420 Ma and ~449 Ma and to the oldest unassignable (i.e., possibly stem or crown) embryophyte spores at ~472 Ma (Clarke et al. 2011, Magallon et al. 2013).

Of the global eukaryotic analyses, the study by Hedges and colleagues (2004) recovered the oldest estimate for the age of the MRCA of stomatophytes: 707 Ma (515–899) (no streptophyte estimate was given). In contrast, Berney and Pawlowski’s (2006) estimate for the embryophytes node was much younger 510 Ma (431 – 645), with stomatophytes dated at ~442 Ma (again, no crown streptophyte age was available). Parfrey et al. (2011) estimated a similar age for stomatophytes (~460 Ma with very small credible intervals) with streptophytes dated at ~745 Ma (~625 – 850).

Although there have been many molecular clock analyses in the last two decades that focus specifically on the streptophytes, here I discuss only two of the most recent studies. Clarke and colleagues (2011) analyzed seven plastid genes and 17 fossil calibrations within the embryophytes using a Bayesian RMC method (MCMCtree, independent rates model). They examined fossil calibration consistency using cross-
validation (Near et al. 2005) as well as extensively explored the impact of uniform versus Cauchy distribution priors on fossil calibrations. Uniform priors have an equal probability that the node’s age is at any point within the given range, while a Cauchy distribution sets maximum probability at point, with decreasing probability for progressively higher or younger dates. Using a uniform prior with hard lower bound and a soft upper bound for all constraints, their ‘best’ estimates for stomatophyte and embryophyte crown nodes were 632 (548 – 750) and 670 Ma (568 – 815), respectively. However, they found that the estimated ages of these deepest nodes were greatly influenced by the upper calibration bound, as well as the nature of the prior distribution (i.e., uniform versus Cauchy) on the fossil bounds. For the above estimates they used a uniform prior with an extremely high upper bound of 1042 Ma for all three nodes (with the lower bound defined by the first fossil evidence for each group).

Magallón et al. (2013) estimated the age of crown stomatophytes as ~458 Ma (446 – 469) and the ancestral embryophyte node at 475 Ma (471 – 480) using the uncorrelated lognormal RMC model implemented in BEAST. While this dataset was comprehensive in terms of taxon sampling (80 taxa) and fossil calibrations (26), only five plastid genes were used. In contrast to the analyses by Clarke and colleagues, Magallón and colleagues set a strong narrow lognormal prior distribution on the age of the MRCA of embryophytes centred at 472 Ma (Magallon et al. 2013), which likely explains the narrow credible region from the posterior distribution they obtained for this node (spanning only 9 Myr).

My taxon sampling does not include the liverworts, the earliest branching lineage within Embryophyta (Magallon et al. 2013). I therefore focus on the age of the common ancestor of living stomatophytes (the split between mosses and all other land plants; Figure 2.1 node e) as a lower bound on the age of embryophytes. I can estimate the upper bound on the age of complex embryophyte-type multicellularity (i.e., diploid multicellular sporophytes) by estimating the age of the basal streptophyte split (Mesostigma versus all other land plants; Figure 2.1, node d). This divergence (i.e., Streptophyta, the upper bracket) was calibrated between 471 Ma (the oldest fossilized embryophyte spores (Rubinstein et al. 2010)), and 600 Ma (following Parfrey and colleagues (2011)). All of my analyses estimated the age of this node to be close to the
upper bound. In fact, the youngest estimates were obtained under the UGam model (~570 Ma) and were associated with very wide confidence intervals spanning more than half the calibration range. In contrast, with LogN and CIR models estimates were older, approaching the upper bound of the calibration, and in one case, just outside of it (LogN + C60, with soft bounds). The lower bracket (i.e., last common ancestor of stomatophytes (Figure 2.1, node e)) shows the opposite trend: CIR and LogN estimates are grouped near the lower calibration bound for that node (425 Ma).

My estimate of ~430 – 450 Ma for Stomatophyta is in agreement with results from Berney and Pawlowski (2006) (~442 Ma), Parfrey et al. (2011) (~460 Ma), and Magallón et al. (2013) (~458 Ma). This congruence is interesting given that these analyses used fairly different age constraints on nodes from this region of the tree. In contrast, Clarke and colleagues (2011) estimated this node to be much older (~630 Ma). However, the results obtained by these authors seem to have been heavily influenced by the root prior, which usually reflects that the data themselves contains little information (Felsenstein 2004, Rannala 2002, Zwickl and Holder 2004).

2.4.5 Red Algae

The first evidence of florideophyte red algae comes from fossils from the Doushantuo Formation dated at 550 – 600 Ma (Xiao et al. 2004). These fossils have been used by a number of molecular clock studies to provide a lower bound on the split of two of the multicellular red algal lineages: the Bangiales and the Florideophyceae (Yoon et al. 2004, Berney and Pawlowski 2006, Parfrey et al. 2011). The second putative red algal fossil often used is the Bangia-like Bangiomorpha pubescens from the Hunting Formation dated at 1198 +/- 24 Ma (Butterfield 2000). As Bangiales and Florideophyceae are a clade, and a number of red algal lineages (some of which are multicellular) branch off before them within the Rhodophyceae (Yoon et al. 2004, Saunders and Hommersand 2004), this fossil is often used as a bound indicating the first appearance of multicellular red algae (Yoon et al. 2004, Berney and Pawlowski 2006, Parfrey et al. 2011), although there has been some argument about its attribution (Cavalier-Smith 2002).

In attempting to date the origin of major eukaryotic photosynthetic groups, Yoon and colleagues (2004) used two rate-smoothing RMC approaches in r8s (Sanderson 2003) to analyze five plastid genes from a large diversity of photosynthetic eukaryote
lineages and seven fossil age constraints (two within the red algae, four within the embryophytes and a maximum age on the root). Because their methods assumed hard bound constraints (and their bounds were narrowly defined around fossil dates), they could only obtain an unconstrained age estimate for the basal red algal split between the bangiophyte+florideophyte clade and the unicellular Cyanidiales. Their estimate for this node was 1,370 Ma (1,350 – 1,416). This age was much older than comparable estimates by Berney and Pawlowski (2006) – they estimated an age of 780 Ma (680 – 950) for this split, while the MRCA of the Floridiophyceae/Bangiales divergence was estimated to be 740 Ma (600 – 929). Note that Berney and Pawlowski did not include the Bangiomorpha calibration in their analyses. Parfrey and colleagues (2011) obtained widely varying estimates for the basal red algal split depending on whether they included the Bangiomorpha and other Proterozoic fossil constraints. Without Proterozoic fossils, they obtained a 95% credible age range for crown red algae of 625 – 959 Ma, whereas, with them, they estimated the range to be 1180 – 1285 Ma. Their estimate for the florideophyte/Bangiales split was ~765 Ma (~630 – 915) when Proterozoic fossil constraints were included, and ~620 Ma (~495 – 700) when excluded.

I estimated the age of the Bangiales/Florideophyceae common ancestor as well as the basal divergence between the unicellular Cyanidiales (Cyanidioschyzon and Galdieria) and the Bangiales/Florideophyceae clade. The Bangiales/Florideophyceae divergence (Figure 2.1, node g) was calibrated by the first appearance of florideophytes in the fossil record (Xiao et al. 2004) with a younger age bound of 550 Ma and an uninformative maximum bound at 3000 Ma. All estimates of the age of this node were within 210 Myr of the lower bound (with soft bounds). It is worth noting that the estimates for this node were likely considerably affected by the estimated age of its immediate ancestral node on our tree. The latter represents the basal divergence in the red algae (Figure 2.1, node f) and was calibrated using the fossil Bangiomorpha pubescens (Butterfield 2000) as a lower bound at 1174 Ma. (Note that while these fossils resemble some Bangiales (Butterfield 2000), here I have treated it as if it were a stem multicellular red algal lineage, following Parfrey et al. 2011). The effect of this fossil is clear when comparing hard and soft bounds: all the hard bound estimates were within 85 Myr of the
minimum calibration, with small confidence intervals constrained by this calibration. However, the soft bound analyses yielded dramatically younger estimates: $\text{LogN} + \text{C60}$ returned an age of 690 Ma, which is 484 Myr younger than the minimum bound set by the fossil; the *Bangiomorpha* fossils seem at odds with other calibrations in my analyses, and seem to be much older than any of the dates estimated from molecular clock data. This discrepancy can, in theory, be explained in a number of ways. For example, either the identification of *Bangiomorpha* fossils as red algae, or the estimated age of the rocks in which they are found, could be in error (although the latter is thought to be unlikely, see discussion in Parfrey et al. (2011) and Knoll et al. (2006)). The discrepancy may also be a result of the stochastic nature of the fossil record. The amount of time between the divergence between the two lineages, and the first recognizable fossil of either of the lineages (i.e.: the fossil used as a minimum bound for that node) is unknown, and could be relatively small or very large. It is possible that many of the fossils used to calibrate minimum bounds in my analyses are significantly younger than the node of interest, because no organism happened to be fossilized, no relevant fossils have been found, or fossils that have been found are not recognizable as belonging to that lineage. By chance *Bangiomorpha* fossilized relatively soon after the divergence of Bangiales/Florideophyceae from other red alga, and its similarity to extant Bangiales allowed it to be classified as a stem multicellular red alga. If this is the case, the older nodes suggested in my hard bound analyses are more accurate, while soft bound estimates underestimate the age of nodes due to the poor fossil record. Alternatively, the changes in the rates of evolution within some lineages, including this one, might be poorly captured by the currently available RMC models.

While my estimates of the most recent common ancestor of Bangiales/Florideophyceae red algae are quite variable (~550 – 720 Ma, with credible intervals spanning up to 500 Myr), they are relatively consistent with those obtained by Berney and Pawlowski (2006) (~700 Ma), and by Parfrey and colleagues (2011) (~620 Ma) when the latter excluded all Proterozoic fossil constraints (including *Bangiomorpha*). However, Parfrey and colleagues’ estimates for this node were much older (~820 Ma) when Proterozoic fossils were included. The basal split between Cyanidiales and the Bangiales/Florideophyceae clade showed much greater differences. My estimates vary
greatly (by up to 500 Myr, with credible intervals even more extreme) depending on the model and style of constraints (Figure 2.3b) and similar degrees of variation in age estimates for this node were found by Parfrey and colleagues (2011) depending on the inclusion/exclusion of Proterozoic fossils constraints. Berney and Pawlowski estimated younger ages (680 – 950 Ma), which were similar to the lower ranges I obtained (e.g., 639 – 771 Ma for LogN+C60) and Parfrey and colleagues (625 – 959 Ma without Proterozoic fossils). The sensitivity of estimates associated with this and the previously discussed node seems to be largely correlated with how ‘strongly’ the Bangiomorpha dates are imposed on it as a constraint. The weaker the constraint (soft bounds in our analysis or removal of the calibration by Parfrey and colleagues), the younger the age estimate for this node.

2.4.6 Relative Ages of Multicellular Groups

According to my analysis, it appears that the Metazoa and Fungi are the oldest groups, with similar age estimates around 700 – 800 Ma when restricted to soft bound analyses (Figure 2.4). The credible intervals on the age of Metazoa are smaller than those for Fungi presumably because this node, as well as many of its descendant nodes, is well constrained by fossil data. It is worth noticing the relatively small difference (~300 – 600 Myr, depending on the RMC model used) between these two nodes and the age estimates for LECA (~1,000 and ~1,350 Ma). This suggests that the first multicellular organisms emerged relatively rapidly after LECA. The Bangiales/Florideophyceae split is estimated to be the next oldest group in soft bound analyses at ~550 – 720 Ma (hard bound analyses yielded substantially older ages because of the fossil Bangiomorpha, as discussed above). Stomatophyte age estimates somewhat younger (430 – 450 Ma) and overlapped with the dictyostelids, which appear to be the youngest multicellular group I considered here (~330 Ma). It should be noted that all of the age estimates (with the exception of the Stomatophyta) are associated with large 95% credible intervals spanning hundreds of millions of years. Thus, despite the large size of my data set, considerable uncertainty in the age estimates persists.
Figure 2.4 Age of the LECA compared to multicellular eukaryotic groups. All estimates were calculated with the LG substitution model and soft bounds. Results are shown for the UGam (triangles), LogN (diamonds), and CIR (crosses) relaxed molecular clock models. Error bars indicate 95% credible intervals. For each node with calibration, the error bar above ‘Cal’ shows the age range used to calibrate the node. Where upper bound is not shown, it was equal to 3000 Ma. M: Metazoa; F: Fungi; D: Dictyostelids; S: Stomatophyta; B: Bangiales/Florideophyceae.

It is important to keep in mind that the dates shown on Figure 2.4 represent age estimates for the most recent common ancestor of each multicellular group, which, without doubt, was not the first multicellular organism in this lineage. The time of first emergence of multicellularity for the lineages I have discussed can be bracketed by considering not only the age of the most recent common ancestor of the group, but also the date of divergence of this lineage from its closest unicellular relative (as I discussed for embryophytes and red algae in the previous section). These bracketed age estimate ranges are given in Table 1; only UGam results are shown as they encompass the range of estimates given by the other two models. Note that the precision of these ‘brackets’ on the age of the emergence of multicellularity depends critically on the taxonomic sampling of the groups. Sparse sampling of the closest unicellular relatives of groups, or the presence of long stem lineages, will artificially inflate the age ranges.
Table 2.2: Limits on the age of emergence of multicellularity in different eukaryotic groups. Lower Limit represents the date of the most recent common ancestor of all sampled members of the multicellular group. The Upper Limit column refers to one node deeper on the rooted tree, i.e., the date of divergence of the group from its most closely related unicellular organism (in parentheses is the node letter(s) from Figure 2.1, where there are two, the second letter represents the upper limit). 95% credible intervals are indicated within the parentheses. Analyses shown were performed using soft bounds, an LG substitution matrix and the UGam relaxed clock model.

<table>
<thead>
<tr>
<th></th>
<th>Lower Limit (Ma)</th>
<th>Upper Limit (Ma)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metazoa (a)</td>
<td>748 (663 - 865)</td>
<td>872 (758 - 1024)</td>
</tr>
<tr>
<td>Fungi (b)</td>
<td>752 (573 - 977)</td>
<td>927 (738 - 1145)</td>
</tr>
<tr>
<td>Dictyostelids (c)</td>
<td>348 (212 - 601)</td>
<td>793 (542 - 1062)</td>
</tr>
<tr>
<td>Embryophytes (e, d)</td>
<td>574 (522 - 604)</td>
<td>852 (654 - 1067)</td>
</tr>
<tr>
<td>Bangiales/Florideophyceae (g, f)</td>
<td>721 (494 - 964)</td>
<td>996 (801 - 1186)</td>
</tr>
</tbody>
</table>

2.5 Conclusions

Here I highlighted some of the progress and difficulties in estimating the age of multicellular eukaryotic lineages using sophisticated ‘relaxed molecular clock’ (RMC) methods. Such estimates are critical if we are to understand how these major evolutionary transitions are correlated with major geological events, such as the oxygen rise in the atmosphere and oceans, or the ‘snowball Earth’ glaciation periods in the Cryogenian. In fact, the association between the origins of animals and oxygen has long been discussed (Lenton et al. 2014, Knoll and Sperling 2014). While the emergence of Metazoa is in broad synchrony with an increase in atmospheric oxygen and the Sturtian glaciation period (~670 – 730 Ma), determining which of these events happened first is not a simple task. It was long thought that a rise in atmospheric oxygen concentrations led to the oxygenation of the ocean, and thus triggered metazoan evolution. The alternative hypothesis would be that the Ediacaran oxygen transition was a consequence rather than cause of animal diversification: surface oceans of dense bacterial populations would have been consumed by filter-feeding animals, while fecal pellets from planktonic bilaterians would have been rapidly sinking from the surface, lessening the oxygen demand (Lenton et al. 2014). To further complicate this issue, there is no evidence for a significant rise in atmospheric or oceanic oxygen levels before the Cryogenian glaciations (including the
Sturtian and Marinoan (~635 – 650 Ma) “Snowball Earth” events), and it was thought that these glaciations posed a major barrier to the survival of eukaryotic life. Yet, the fossil record indicates that numerous eukaryote lineages continued through the glaciations. In fact, my results (and those of others I discuss) postulate that the last common ancestor of metazoans and Fungi existed ~700 – 800 Ma, possibly before the Cryogenian glaciations. Indeed, it has been argued that sponges were present during, and perhaps, even before these glaciations (Love et al. 2009). If confirmed, this would suggest that the origins of the first metazoans could not have occurred as a response to increasing ocean oxygenation.

This example emphasizes the importance of being able to pinpoint precisely the timing of the origin of multicellular lineages and correlate them with respect to the ancient geochemical record. However, my results clearly show that the obtained molecular dating estimates depend heavily on the models and methods used, and on the nature and treatment of fossil calibrations. This, combined with the significant uncertainty associated with most of the age estimates (often spanning many hundreds of millions of years) suggests that my conclusions remain quite tentative. Hopefully, as further investigations fill out the Proterozoic fossil record associated with protistan eukaryotes, more genomic data is gathered from diverse eukaryote lineages and improvements are made in relaxed molecular clock modeling and methods, we will obtain more precise estimates of the ages of multicellular groups.

2.6 Acknowledgements

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Chapter 3: Integrins and the Evolution of Animal Multicellularity


3.1 Introduction

The evolution of cell adhesion was an important step in the transition from unicellular organisms to multicellular organism, including animals. New cell adhesion genes present in multicellular groups encode adhesion proteins, or change cell division/cell wall formation so that cells remain attached after division (Bonner 2000, Knoll 2011). Animals have specialized adhesion proteins, which are different from adhesion strategies found in other multicellular groups (Abedin and King 2010). One important class of animal cell adhesion proteins are the integrins: heterodimeric transmembrane receptors involved in cell signaling and adhesion to the extracellular matrix (Hynes 2002).

Early research on cell adhesion in vertebrates identified areas near the cell membrane where fibrils of actin end together as a bundle, near points of contact between the cell and the extracellular matrix (ECM) (Horwitz 2012). The molecule that connected the outside ECM to the internal cytoskeleton remained unknown until 1986, with the isolation of a cDNA clone which was sequenced for structural analysis (Tamkun et al. 1986). The protein contained a relatively large N-terminal segment outside the cell (including three cysteine rich repeats), a transmembrane region, and a short cytoplasmic region. This protein was termed ‘integrin’ for its integral role in connecting cells to extracellular matrix (Tamkun et al. 1986). The discovery of this β subunit was followed by the characterization of the α subunit, and soon a number of integrins were recognized as a group of homologous cell receptors involved in cell adhesion, migration, thrombosis and immune system function (Hynes 1987).
Structural studies of integrins (reviewed in Ginsberg (2014) and Fu et al. (2012)) show that both the α and β subunit have the general structure of a globular head domain, a long stalk region, a single short transmembrane region and a short cytoplasmic tail. Despite the superficial similarities in their structures, the two subunits are not observably homologous, and differ in their domain compositions. In integrin β (ITB), the head domain is made up of a von-Willebrand Factor A domain (Whittaker and Hynes 2002) and the stalk is made up of 3 - 4 repeating EGF units, a domain held together by disulphide bonds, some of which are formed by a conserved cysteine-rich motif. The intracellular portion contains two NPXY motifs, which can be phosphorylated to modulate interactions with other proteins (Anthis and Campbell 2011). ITB4 is an exception: it has intracellular Calx-β domains which interact with intermediate filaments instead of actin. Integrin α (ITA) has a head region made up of a seven-bladed β propeller domain, each containing an FG-GAP motif. The stalk is made up of a thigh domain and two calf domains, separated by a flexible knee region that is important for integrin conformational regulation (Fu et al. 2012). The integrin ligand-binding site is formed at the interface between the α and β subunits, and generally binds extracellular matrix proteins, although some integrins have been co-opted for cell-to-cell communication in the immune system (Humphries et al. 2006). Mammals have 18 α subunits and 8 β subunits which combine to form 24 distinct pairs, with various functions in immune response, cell adhesion, and cell signaling (Hynes 2002).

Integrins can exist in two conformations: a low-affinity conformation where the stalk region is bent, causing the ligand-binding site to point towards the cell membrane, and an active conformation where the stalks are straight, exposing the ligand-binding site (Zhu et al. 2008, Xiong et al. 2002, Vicente-Manzanares et al. 2009). This conformational change can be triggered by talin, which binds to the cytoplasmic region of integrin β, or by ligand binding. As a result, integrins are important to both outside-in and inside-out cell signaling (Anthis and Campbell 2011). Binding of integrins to the ECM results in upregulation of proliferation genes, and both kinds of signaling may be modulated by mechanical stress (Dufort et al. 2011).

Integrins act in cell migration or adhesion by connecting to the actin cytoskeleton through a number of scaffolding proteins (Anthis and Campbell 2011, Vicente-
Manzanares et al. 2009). Talin binds the ITB cytoplasmic region with an N-terminal FERM domain, while the rod shaped C-terminal region contains several places where α-actinin, an actin-binding protein, can bind. ILK (integrin linked-kinase), PINCH and paxillin make up the IPP complex, which serves as a scaffold for protein-protein interaction and modulates signaling. Focal adhesion kinase (FAK) and c-Src (a proto-oncogene and kinase) are involved in downstream signaling, promoting proliferation when cells are bound to the ECM. Disruption of any of the genes that associate with them can lead to many diseases, in particular cancer (Winograd-Katz et al. 2014).

Cell migration is important for development and immune system response, two important functions of multicellular organisms. Integrins provide a stable connection to the ECM that is necessary for cells to be able to move along it (Vicente-Manzanares et al. 2009). As the leading edge of the cell protrudes, new focal adhesions form when integrins bind the ECM. Actin polymerization farther forward then pushes the edge of the cell forward. At the trailing edge of the cell, focal adhesions disassemble, releasing the cell and allowing it to move forward. Scaffolding proteins (for example, talin) work as a molecular clutch to transfer traction from the integrins to movement powered by actin polymerization.

Integrins are not present in other multicellular groups such as plants and fungi. Choanoflagellates, the closest unicellular relative to metazoans, do not have integrins either, which initially led to the assumption that integrins were metazoan specific. Recently, integrins have been discovered in the holozoan Capsaspora owczarzaki and the apusomonad Thecamonas trahens (Sebé-Pedrós et al. 2010, Sebé-Pedrós and Ruiz-Trillo 2010). These findings suggest that integrins evolved not during the origin of metazoa, but earlier, nearer to the divergence of Amoebozoa and other Unikonts, with a secondary loss in Fungi (Figure 3.1). The function of integrins in protists is currently unknown, but could be related to motility or predation. The relationship between Fungi and Animals, which together form the Opisthokonts, has long been known (Wainright et al. 1993) but more recently many newly discovered or misplaced unicellular organisms have been included in this group (Mendoza et al. 2002). Elucidation of the phylogenetic relationships within this group is ongoing (Torruella et al. 2015), and these organisms can
be valuable for determining the genetic changes necessary for the transition to a multicellular lifestyle.

The Breviatea are an anaerobic protistan group recently found to be more closely related to Opisthokonts than to the other supergroups. The discovery and transcriptome sequencing of the breviate *Pygsuia biforma* provided enough data to confidently assign the breviates to a position sister to the Opisthokonts + Apusomonad clade (Brown et al. 2013). *Pygsuia biforma* is an anaerobe which lives as two distinct cell types: swimming cells and adherent cells. During logarithmic growth in culture, swimming cells with two long flagella predominate, while in older cultures adherent cells with one long flagellum and one short flagellum are more common. The adherent cell crawls along the surface using many filose pseudopodia, which can extend behind it leaving a long trail (Brown et al. 2013).

The newly discovered position of breviates as a sister lineage to the Opisthokonta prompted a search in the transcriptome of *Pygsuia biforma* for integrins and related proteins. I discovered homologs of an integrin α and an integrin β subunit, both with conserved motifs required for the functions of metazoan integrins. *Pygsuia* also expresses ILK, PINCH, paxillin, talin and α-actinin (Figure 3.1). To aid further work on elucidating the functions of these proteins in protists, I obtained and tested antibodies against the head domains of *Pygsuia* integrin α and β, the cytoplasmic region of integrin β, and the FERM domain of talin.

### 3.2 Methods

#### 3.2.1 Homology Searches

Bilaterian integrin-mediated adhesion complex genes were obtained from the Homologene database (http://www.ncbi.nlm.nih.gov/homologene/). Sequences from each Homologene group were aligned in MAFFT using the LINS-i option (Katoh and Toh 2008) Hidden Markov Model (HMM) profiles were created from the alignments using HMMbuild of HMMER 3.0 (http://hmmer.org/). A six frame translation of the *Pygsuia biforma* transcriptome and protein datasets from genomes of non-bilaterian animals and other representative eukaryotes were searched for homologs using HMMsearch with a full sequence threshold of $1e^{-5}$ (see (Brown et al. 2013) for details on transcriptome
culture conditions, library preparation, sequencing and assembly). Domains of the HMMsearch hits were identified using the HMMpfam and SMART databases of InterProScan (Zdobnov and Apweiler 2001). The putative homologs found by HMMSEARCH were then screened using a reciprocal Basic Local Alignment Search Tool (BLAST) technique; BLASTP of the putative ortholog against the National Center for Biotechnology Information (NCBI)’s non-redundant protein database. Criteria for homologs of each protein were then established: for some proteins, domain architecture with minimal changes (e.g. possibly a domain missing, or different numbers of repetitive domains) was used as an inclusion criterion; for others, domain architecture was considered uninformative, but blast confirmed similarity to the protein of interest. HMM searches for particular diagnostic domains (e.g: SH2 [PFAM PF00017] for SRC and Focal_AT [PFAM PF03623] for FAK) were also performed on the transcriptome and proteomes.

3.2.2 Pygsuia biforma cDNA Preparation

Pygsuia biforma was grown in a 250 mL tissue culture flask previously seeded with Klebsiella pneumoniae in Sonneborn’s Paramecium medium (ATCC medium 802) made with filtered autoclaved sea water for five days with limited headspace, and then collected by centrifugation. 10 mL of TRIzol® (Life Technologies) was added, and the mixture was incubated for 15 minutes at 4°C. 2 mL of chloroform was added and the organic and aqueous layers were separated by centrifugation at 3220 g in an Eppendorf 5810R centrifuge for 75 minutes at 4°C. The aqueous layer was collected, and 5 mL of isopropanol was added. The sample was centrifuged at 3220 g for 45 min 4°C, and the supernatant was discarded. The pellet was washed with 75% ethanol, centrifuged for 10 minutes at 3220 g (4°C) and dried for 15 minutes at room temperature. The pellet was then resuspended in nuclease-free water and a portion was resolved by agarose gel electrophoresis to assess quality. The remaining RNA was precipitated with NaCl in ethanol and stored at -80°C. Eukaryotic mRNA was selected from the total RNA sample using the Poly A Purist MAG kit (Ambion Inc, TX), and reverse transcribed using the SMART MMLV reverse transcriptase (Clontech Laboratories, Inc.) using poly deoxythymidine as a primer.
3.2.3 Pygsuia Whole Cell Lysates

Pygsuia was grown in 15 mL conical flasks seeded with a clump of Klebsiella pneumoniae and filled with culture medium (see above) with less than 1 mL of headspace. Cultures were grown for 15 days and then centrifuged at 400 g and 4°C for 20 minutes, and all but four mL of media was removed. This sample was then centrifuged for 10 minutes at 800 g and for another 10 minutes at 3220 g. All supernatant was removed, and approximately 100 μL of 10 mM phosphate buffered saline (PBS) was added. Cells were mechanically disrupted by syringing, and the whole cell lysate was tested for antibody reactivity along with the recombinant protein samples, described below.

3.2.4 Expression of Recombinant Protein

DNA encoding Pygsuia integrin complex protein domains was amplified from Pygsuia biforma cDNA (for ITB, both cytoplasmic and ‘head’ domains) or synthesized by GeneArt (Life Technologies – for talin (FERM domain) and ITA (head domain)). TA cloning was used to insert PCR products and synthesized DNA into the pGEM–T Easy vector, and restriction enzymes were used to insert the protein-coding DNA into expression plasmids. pET-16b (Novagen) was used for expression of ITB head, ITA head, and talin FERM domains, while the cytoplasmic region of ITB was expressed in pET-32 a + (Novagen), which expresses the insert as a thioredoxin fusion protein. Proteins were expressed in BL21(λDE3) (ITB cyto, ITA head), BL21(λDE3) pLys (ITB head) or C43 (talin FERM) cells, with induction by 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG). Cells were mechanically disrupted by syringing, and sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) analysis of the whole cell lysate with Coomassie staining confirmed the expression of protein. For the cytoplasmic region of ITB, His-tagged recombinant protein was purified using Clontech’s TALON® Metal Affinity Resin.

3.2.5 Antibody Testing on Recombinant Protein

Polyclonal rabbit antibodies were raised against peptides from Pygsuia ITA head domain, ITB cytoplasmic region, ITB head domain and talin FERM domain by GenScript Inc. (USA). For each target protein, immunogenic peptides that were 13 amino acids in length were identified by GenScript, and one from each domain of interest was chosen.
based on its degree of sequence conservation with protist homologues (i.e. highly conserved peptides were selected, Table 3.1).

**Table 3.1 Peptides used to raise antibodies.** Cysteines (C) in bold were conjugated to the peptide to aid in antibody production, and are not present in the *Pygusia* sequence.

<table>
<thead>
<tr>
<th>Protein</th>
<th>Domain</th>
<th>Position</th>
<th>Peptide</th>
</tr>
</thead>
<tbody>
<tr>
<td>ITB</td>
<td>cytoplasmic</td>
<td>1776</td>
<td>CNALYEHGNTQADGSQ</td>
</tr>
<tr>
<td>ITB</td>
<td>head</td>
<td>117</td>
<td>VLEDLSGSFGDDLRC</td>
</tr>
<tr>
<td>ITA</td>
<td>head</td>
<td>555</td>
<td>SSIPSEIPGPSTSSC</td>
</tr>
<tr>
<td>Talin</td>
<td>FERM</td>
<td>86</td>
<td>KIRPLRLKLSDETTC</td>
</tr>
</tbody>
</table>

His-purified recombinant protein or whole cell lysates of *E. coli* expressing recombinant protein were tested for antibody reactivity by western blotting. After separation by SDS-PAGE, samples were transferred to a polyvinylidene difluoride (PVDF) membrane using BioRad’s Turboblot semi-dry blotting system. Membranes were blocked using 5% skim milk powder in Tris buffered saline (TBS) containing 0.1% Tween 20 (TBS-T), and then incubated with primary antibody in 5% milk in TBS-T overnight at 4°C (1:500 for ITA head antibody, 1:10,000 for other antibodies). After washes in TBS-T, blots were incubated with 1:50,000 horseradish peroxidase (HRP)-conjugated anti-rabbit IgG secondary antibody (Sigma) for one hour at room temperature. After further washes with TBS-T, blots were developed with electrochemiluminescence (ECL) reagents (Amersham), and photographed using a FluorChem E imaging system (Protein Simple). When whole cell lysates were used, cells transformed with an empty expression plasmid were used as negative controls.

### 3.3 Results

*Pygusia biforma* expresses a complement of integrin mediated adhesion proteins similar to that of the apusomonad *Thecamonas trahens* (Fig 3.1). An α integrin, a β integrin, and many of the associated scaffolding proteins were identified in the transcriptome data. As in *Thecamonas trahens*, neither FAK nor c-Src were present,
Figure 3.1: Distribution of integrin adhesion machinery in eukaryotes. The integrins and related proteins in Pygsuia biforma and Sphaeroforma arctica (red boxes) were discovered in my analysis, although the presence of integrins in Sphaeroforma has been previously acknowledged (Sebé-Pedrós and Ruiz-Trillo 2010). Numbers indicate number of integrin subunits, filled circles indicate presence of the gene and empty circles indicate putative or degenerate homologs. Adapted from Sebé-Pedrós et al. 2010 consistent with classifying these proteins as holozoan innovations. An ITB and associated integrin scaffolding and signaling proteins were also found in the icthyosporean Sphaeroforma arctica, but no ITA was found, so it is unclear how its integrin receptor would function. While these proteins were recognizable homologs to the integrin-mediated adhesion machinery in animals, they did show some interesting differences in domain composition (Figures 3.2, 3.3, 3.4).

Pygsuia biforma has one integrin α and one integrin β. Both subunits are significantly longer than their animal homologs. In each case, the expansion comes from the stalk region of the protein. Animal β integrins contain three to four EGF, while Capsaspora has four to seven, and Thecamonas has seven. The Sphaeroforma arctica ITB is of a similar length to the ITBs of these organisms, with three EGF repeats. The ITB stalk is much longer in Pygsuia biforma, which has 29 EGF repeats (Figure 3.3), as well as 27 repeats of the CXCXXXCXC motif, which in model system homologs form disulphide bonds to maintain the structure of the domain (Table 3.2). The stalk region of
**Homo sapiens** 801 AA

Figure 3.2 Domain architectures of protistan integrin β proteins. SP, signal peptide (SIGNALP-NN(euk)); TM, transmembrane domain; ITB head, integrin b chain (PF00362); EGF, epidermal growth factor; extracellular (IPR013111); BT, integrin-B tail (PF07965); BC, integrin-B cytoplasmic region (PF08725). Vertical purple lines represent cysteine-rich repeats (CxCxxCxC) (PS00243). The numbers of EGF and CxCxxCxC repeats are indicated. Adapted from Brown et al. 2013.
Figure 3.3 Domain architectures of protistan integrin α proteins. SP: signal peptide; ITA head; FG: FG-GAP repeats; TM: transmembrane region. ITA head, integrin α, β-propeller (IPR013519); FG, FG-gap (PS51470); ITA leg, integrin α-2 (IPR013649); TM, transmembrane domain. Adapted from Brown et al. 2013.
ITA is even more dramatically extended, resulting in a protein that is three times as long as its metazoan orthologs (Figure 3.2). Animal ITAs contain an ITA leg domain (integrin a-2 - IPR013649), which has been well studied for its effects on the conformational regulation of integrins (Xiong et al. 2001, Luo et al. 2007). This domain is not present in previously described protistan integrins (Sebé-Pedrós et al. 2010), or the ITAs described here. It is therefore unclear whether protistan integrins have the same kinds of regulatory properties.

**Homo sapiens** 2542 AA

**Capsaspora owczarzaki** 2508 AA

**Thecamonas trahens** 3776 AA

**Pygsuia biforma** 2451 AA

![Diagram]

Figure 3.4 Domain architectures of protistan talins. *Homo sapiens* talin-2 (GENBANK NP_055874.2, which is typical of metazoan talins), and *Thecamonas trahens* (updated version of the talin ortholog noted in Sebé-Pedrós et al. 2010, from Broad Institute locus AMSG_12193.2). Abbreviations: FERM (Pfam: PF09379) (FERM N), PF00373 (FERM M), PF02174 (I = IRS) TM= Talin middle (Pfam: PF09141), VBS = Vinculin binding site (Pfam: PF08913), I/LWEQ (actin binding domain, Pfam: PF01608), P= Paired amphipathic helix repeat (Pfam: PF02671), H= Histone deacetylase interacting (Pfam: PF08295), V= Villin headpiece (Pfam: PF02209) Adapted from Brown et al. 2013.

Functional motifs are conserved in the ITBs of *Pygsuia* and *Sphaeroforma* (Table 3.2). The metal ion-dependent adhesion site (MIDAS) is perfectly conserved in both protists, while the ligand-associated metal binding site (LIMBS) and adjacent to MIDAS
(AMIDAS) sites are recognizable with only one amino acid change. In animal integrins MIDAS coordinates a divalent cation with an oxygen from the aspartate (D) of the RGD motif in the ligand (Xiong et al. 2002), while the AMIDAS site binds another divalent cation nearby. LIMBS coordinates a divalent cation when ligand is bound, stabilizing the area of the protein that interacts with the ligand (Luo et al. 2007). The conservation of these sites in *Pygsuia* and *Sphaeroforma* as well as in *Thecamonas* and *Capasaspora* (Sebé-Pedrós et al. 2010) suggests that protist integrins may bind ligands in a manner similar to that of animal integrins, perhaps ones containing an RGD motif. The presence of an NPXY motif in the cytoplasmic region of the ITBs found in *Pygsuia* and *Sphaeroforma* suggests that interactions between the integrin receptor and other proteins such as talin may be conserved between animals and protists.

### Table 3.2 Conserved motifs in integrin β

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<th>Ion binding motifs</th>
<th>Metazoa</th>
<th>Sphaeroforma arctica</th>
<th>Pygsuia biforma</th>
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<tr>
<td>MIDAS</td>
<td>Amino acid (AA)</td>
<td>Position</td>
<td>AA</td>
</tr>
<tr>
<td>D</td>
<td>D</td>
<td>119</td>
<td>D</td>
</tr>
<tr>
<td>S</td>
<td>S</td>
<td>121</td>
<td>S</td>
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<tr>
<td>D</td>
<td>D</td>
<td>123</td>
<td>D</td>
</tr>
<tr>
<td>E</td>
<td>E</td>
<td>232</td>
<td>E</td>
</tr>
<tr>
<td>D</td>
<td>D</td>
<td>260</td>
<td>D</td>
</tr>
<tr>
<td>AMIDAS</td>
<td>S</td>
<td>123</td>
<td>S</td>
</tr>
<tr>
<td>D</td>
<td>D</td>
<td>126</td>
<td>D</td>
</tr>
<tr>
<td>D</td>
<td>D</td>
<td>127</td>
<td>D</td>
</tr>
<tr>
<td>D</td>
<td>D</td>
<td>260</td>
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</tr>
<tr>
<td>D</td>
<td>A</td>
<td>389</td>
<td>V</td>
</tr>
<tr>
<td>LIMBS</td>
<td>D/E</td>
<td>167</td>
<td>D</td>
</tr>
<tr>
<td>ligand-associated metal binding site</td>
<td>N</td>
<td>227</td>
<td>G</td>
</tr>
<tr>
<td>D</td>
<td>D</td>
<td>229</td>
<td>D</td>
</tr>
<tr>
<td>P</td>
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</tr>
<tr>
<td>E</td>
<td>E</td>
<td>232</td>
<td>E</td>
</tr>
<tr>
<td>Repeated motifs</td>
<td>Number of repeats</td>
<td></td>
<td></td>
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<tr>
<td>CXCXXCXCX</td>
<td>3-4</td>
<td>5</td>
<td>27</td>
</tr>
<tr>
<td>NPXY</td>
<td>1-2</td>
<td>1</td>
<td>1</td>
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Many cytoplasmic integrin–related scaffolding and signaling proteins are expressed in *Pygsuia biforma*. Homologs of the scaffolding proteins PINCH, paxillin, talin and α-actinin were present, as well as integrin linked kinase (ILK). FAK and c-Src were not present, as in *Thecamonas trahens* (Sebé-Pedrós et al. 2010), suggesting that these are holozoan-specific proteins. More surprising is the apparent absence of the actin binding proteins vinculin and parvin. Consistent with the lack of a vinculin homolog in its RNAseq data, *Pygsuia*’s homolog of talin lacks a vinculin-binding domain (Figure 3.4), which is present in metazoan and other protist talins.

**Figure 3.5** Antibodies raised against peptides are reactive to *Pygsuia* proteins expressed in *E. coli*. Left: Coomassie stained membranes, Right: western blots; chemiluminescent signal corresponding to antibody binding appears white, while red shows the outline of the membrane. L: ladder; -: whole cell lysates of *E. coli* expressing *Pygsuia* recombinant protein (except for panel a – here recombinant protein was isolated from whole cell lysates by affinity purification using the added His-tag), - : whole cell lysates from *E. coli* with empty plasmid, P: *Pygsuia* whole cell lysates.
Antibodies were raised against peptides from *Pygsuia* integrin α (head domain), integrin β (head domain and cytoplasmic region) and talin (FERM domain) (Table 3.1). All four antibodies were reactive against recombinant protein constructs expressed in *E. coli* (Fig 3.5), with little cross-reaction observed with other proteins. ITB cytoplasmic region was purified using the His-tag on the recombinant protein, producing a relatively pure band of recombinant protein in the positive control lane (Fig 3.5 a). For the other three protein domains the chemiluminescent signal corresponds to the one band present in the positive control lane, but not the negative control (expression with empty plasmid) lane (Fig 3.5 b – d). For reasons yet unclear (see Discussion), tests of both the ITB head and Talin FERM domain antibodies by western blots of whole cell lysates of *Pygsuia* cultures showed no detectable signal (Fig 3.5 b and d, right-most lane).

3.4 Discussion

The discovery of integrins and related proteins in *Pygsuia biforma* is evidence for an earlier emergence of the integrin mediated adhesion complex (IMAC) than previously thought (Sebé-Pedrós et al. 2010) (Fig. 3.1). With the results presented here, one can recreate a more complete history of the evolution of the IMAC through eukaryotic evolution. α-actinin is ubiquitous within eukaryotes as an actin binding protein (Fig. 3.1). PINCH, paxillin and talin are amorphean proteins, as they are present in both Amoebozoa and Obazoa. In *Dictyostelium*, an aggregative multicellular organism, talin has been shown to interact with non-integrin cell adhesion proteins (Cornillon et al. 2008). Integrin α, integrin β and ILK are obazoan innovations, as confirmed by the description of IMAC proteins in *Pygsuia*. *Pygsuia* lost both vinculin and the vinculin binding site in talin, which are both present in many other amorphans. Parvin is an innovation of the Apusomonads + Opisthokonts. Within the opisthokonts, IMAC proteins were reduced in Fungi, and enriched in Holozoa. No Fungi have integrin α or integrin β, but the presence of associated signalling proteins in basal fungi (*Spizellomyces, Batrachochytrium* and *Allomyces*; Fig 3.1) suggests a gradual loss of IMAC proteins in this lineage. Holozoa enriched the complement of IMAC proteins by adding the kinases FAK and c-Src, as well as by increasing the numbers of integrin subunits (four integrin αs and four integrin βs in *Capsaspora*, and up to eighteen integrin αs and eight integrin βs in mammals). The
IMAC is an obazoan innovation that has acquired particular importance in Metazoa for its role cell adhesion.

In animals, integrins bind proteins in the extracellular matrix (ECM). As Pygsuia is a unicellular organism, it is unclear what its integrins are binding. The same was true of Capsaspora until the discovery of a multicellular aggregative stage in that organism’s life cycle (Sebé-Pedrós et al. 2013b). Aggregation and the secretion of an ECM was observed, after which cells became cysts. Upregulation of integrins, integrin-associated proteins, tyrosine kinases and a laminin G domain containing protein were observed during aggregation. A currently unobserved multicellular life cycle stage of Pygsuia could explain the presence of integrins. Alternatively, integrins may function in interactions between Pygsuia cells during sexual reproduction, where the integrin ligand is part of a surface protein on another cell.

Another possibility for the function of integrins is motility. Animal cells use dynamic integrin binding to attach to and move across the ECM (Vicente-Manzanares et al. 2009). Thus Pygsuia integrins could bind substrate, creating the connection between the environment and the actin cytoskeleton that is needed to push the cell forward. Adherent Pygsuia cells move by extruding multiple filose pseudopodia, which remain attached to the substrate and move along the ventral side of the cell in a conveyor belt fashion (Brown et al. 2013) (this is similar to movement seen in another breviate, Breviata anathema (Heiss et al. 2013)). The connections to the substrate observed in moving Pygsuia may be mediated by integrins.

Another alternative is that Pygsuia uses integrins to interact with its bacterial prey. Pathogens, both bacteria and viruses (Hussein et al. 2015), exploit animal integrins by using them to adhere to host cells, or to induce phagocytosis. For example, the first steps of Yersinia infection of host cells involves adhesion with the aid of a protein that binds to integrins with a higher affinity than does fibronectin (Mikula et al. 2012). Mutualistic integrin-mediated interactions between bacteria and Pygsuia are also possible. Interactions between prokaryotes and protists that benefit both organisms have been observed, for example in the associations of nitrogen-fixing bacteria with protists (Kneip et al. 2007). Integrins could facilitate such interactions by adhering to bacteria, keeping them in place on the surface of Pygsuia as symbionts.
Localization of integrins and related proteins within *Pygsuia* cells could distinguish among these possible functions. I have obtained antibodies for four domains of three proteins important to the function of integrin-mediated adhesion in animals, and shown that they are reactive in western blots of recombinant protein. The antibodies did not react with *Pygsuia* whole cell lysates, likely because of low density of *Pygsuia*, although it is also possible that these proteins are not expressed under the culture conditions. As *Pygsuia* feeds on bacteria, and bacteria greatly outnumber *Pygsuia* in culture, there is much more bacterial protein compared to protistan protein in whole cell lysate, and large culture volumes may be necessary to obtain a positive western signal. The peptides used to raise the ITB and talin antibodies are similar to the corresponding regions in *Thecamonas* and *Capsaspora* orthologs of these proteins, so the antibodies I developed may be useful for elucidated the function of integrins in these organisms.

### 3.5 Conclusion

*Pygsuia biforma*, a newly described breviate, expresses both subunits of the animal cell adhesion receptor integrin. It also expresses several proteins (ILK, paxillin, talin and α-actinin) that in animals are associated with integrins and help make the connections between the ECM outside the cell and the actin cytoskeleton within the cell. The functions of these proteins in *Pygsuia* and other protists remain unknown, but could be related to motility, predation or other interactions with bacteria, or with a sexual stage of the life cycle. As tools to study these proteins further, I designed antibodies to cross-react with conserved peptides from the head domains of *Pygsuia* ITA and ITB, the cytoplasmic region of ITB, and the FERM domain of talin. Polyclonal affinity-purified antibodies were raised against these peptides and confirmed to be reactive with recombinant proteins in western blots. In summary, I show that integrins evolved earlier than previously thought, and that the newly described breviate *Pygsuia biforma* may provide insights into the genetic toolkit related to animal multicellularity.
Chapter 4: Final Conclusion

There are many independent origins of multicellularity in eukaryotes. Multicellular organisms can form by aggregation of cells or through division of cells that subsequently stick together; such multicellular forms account for the majority of the macroscopic biosphere. As transitions to multicellularity in many groups occurred millions of years ago, the steps involved in these transitions are difficult to study. However, comparisons between various multicellular groups, and between multicellular organisms and their unicellular relatives, help us understand how, when and why multicellularity evolved. The evolutionary transition from unicellular to multicellular has been replicated in several experiments (Boraas et al. 1998, Ratcliff et al. 2012, Ratcliff et al. 2013), demonstrating that selection pressures such as predation can precipitate the transition. Multicellularity involves a switch of the level of selection, meaning that evolutionarily favoured traits might benefit the organism at the expense of the individual cell (Grosberg and Strathmann 2007). To form a successful multicellular organism cells must adhere to and communicate with one another. Using comparative genomics, one can identify the necessary genetic changes that result from the aforementioned selection pressures to gain an understanding of the mechanisms underpinning unicell-to-multicell transitions.

Reconstructing the evolutionary transitions from unicells to multicells requires knowledge of the relationships between extant unicellular and multicellular organisms. Comparisons that are made without consideration of the organismal relationships (phylogeny) can often be misleading. For example, studies that compare model animal systems to unicellular yeast (e.g. Nam et al. 2015, Zaidel-Bar 2009), are comparing a multicell to a unicell that secondarily evolved from a multicellular lineage (Fungi), and underwent many changes to adapt to a saprotrophic lifestyle. While these comparisons benefit from the large amount of data available from model systems, they suffer from the lack of consideration of unicellular amorpheans, in particular unicellular holozoans, which are more closely related to animals than is yeast. As demonstrated by the discovery of integrins in Pygsuia and other protists, some aspects of animal cell biology may be more similar to those of organisms outside Opisthokonta than those of the more closely related Fungi. Increased study of protistan opisthokonts (e.g. choanoflagellates,
filasterians, ichyosporeans, apusomonads and breviates) has transformed how we think of animal origins. Genes that control metazoan-specific processes, such as the development of metazoan body plans, have now been found in unicellular organisms, changing how we understand the genetic transition from unicellularity to multicellularity.

While relationships between multicellular groups (and within eukaryotes as a whole) are becoming better resolved (Parfrey et al. 2010, Brown et al. 2012a, Burki et al. 2007), the time in geological history when each group diverged from its unicellular sister taxa is less clear. As fossils are rare for any species, and in particular for unicellular organisms, there are few fossils that can be used to calibrate ancient nodes in molecular clock analyses. Relaxed molecular clock methods seek to answer these questions by using paleontological evidence to calibrate points on phylogenetic trees, and providing estimates of ages on the remainder of nodes on the tree based on fitted models of evolutionary rate change over time. It is not clear which of these models of rate evolution are the most accurate, so a large range of dates can be estimated for any particular divergence. Some of this uncertainty could be remedied with more data (new fossil calibrations and more sequence data from under-sampled lineages), but the inherent uncertainty in fossil dates and the complexity of modeling the variation of evolutionary rates over time means that most age estimates will continue to be associated with relatively large uncertainty.

While each eukaryotic multicellular group is unique, and offers its own interesting biological questions, there is much to be gained from comparisons between distantly related organisms. Comparisons between groups that evolved multicellularity independently help pinpoint commonalities in the conditions that led to the emergence of multicellularity. Conversely, they illustrate the contingencies present in evolution, as each multicellular group has its own unique features. Estimates of the timing of the emergence of various multicellular groups add to our understanding of eukaryotic relationships, allowing correlations of the presence of particular types of organisms with other geological data, which provides a better understanding of ancient ecologies. Comparisons between multicellular groups and their unicellular relatives allow the investigation of how cells adapt to survive as a part of a larger individual, and what original features were co-opted from the unicellular life to facilitate the multicellular
lifestyle. By understanding the relationships between multicellular groups and their unicellular relatives, one is able to piece together the history of the transition from unicellularity to multicellularity, revealing the mechanics of an important step in the evolution of life on Earth.
References


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