Why It Is Time to Look Beyond Algal Genes in Photosynthetic Slugs

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Abstract

Eukaryotic organelles depend on nuclear genes to perpetuate their biochemical integrity. This is true for mitochondria in all eukaryotes and plastids in plants and algae. Then how do kleptoplasts, plastids that are sequestered by some sacoglossan sea slugs, survive in the animals’ digestive gland cells in the absence of the algal nucleus encoding the vast majority of organellar proteins? For almost two decades, lateral gene transfer (LGT) from algae to slugs appeared to offer a solution, but RNA-seq analysis, later supported by genome sequencing of slug DNA, failed to find any evidence for such LGT events. Yet, isolated reports continue to be published and are readily discussed by the popular press and social media, making the data on LGT and its support for kleptoplast longevity appear controversial. However, when we take a sober look at the methods used, we realize that caution is warranted in how the results are interpreted. There is no evidence that the evolution of kleptoplasty in sea slugs involves LGT events. Based on what we know about photosystem maintenance in embryophyte plastids, we assume kleptoplasts depend on nuclear genes. However, studies have shown that some isolated algal plastids are, by nature, more robust than those of land plants. The evolution of kleptoplasty in green sea slugs involves many promising and unexplored phenomena, but there is no evidence that any of these require the expression of slug genes of algal origin.

Key words: lateral gene transfer, kleptoplasty, photosynthesis, plastid biology, photosynthetic sea slugs.

Introduction

Sacoglossa are considered one of nature’s curiosities. Inside some of these sea slugs, plastids sequestered from algae can continue to photosynthesize for weeks, or even months, in the absence of algal nuclei (Greene 1970; Rumpho et al. 2001; Händeler et al. 2009). That is conspicuous, because when land plant plastids are isolated and removed from their cellular context they rapidly degrade (Leegood and Walker 1983; Seftor and Jensen 1986; Polanská et al. 2004; Green et al. 2005). With the description of endosymbiotic gene transfer (EGT; Martin et al. 1993) and the concomitant genome reduction the organelles experienced (Timmis et al. 2004), the prime cause for the instability of isolated plastids quickly became apparent: the majority of proteins working in plastids are nuclear-encoded and posttranslationally imported from the cytosol (McFadden 2014). Hence, the duration with which kleptoplasts are kept functional in animal cells in the absence of algal nuclei encoding a 1,000+ plastid proteins presents an obvious contradiction. This required an explanation and in 1996 (Pierce et al. 1996) it was proposed that slugs had...
acquired algal genes that encode proteins servicing the plastids through lateral gene transfer (LGT). Once the idea was presented, it was destined to be tested.

**There Is No Evidence for Lateral Transfer of Algal Genes in Slugs**

Let us first take a look at what we should expect if genes of algal origin were to play a role in kleptoplast maintenance. The slugs are sometimes referred to as “crawling leaves,” because their entire appearance of the species in question (e.g., *Elysia chlorotica*, *Elysia timida*, and *Elysia viridis*) is greenish and in a few cases indeed leaf-like. Scientists noticed this already more than 150 years ago. They were particularly intrigued by the digestive tubules that pervade almost the entire body, sometimes including the head, and whose cells house the kleptoplasts (fig. 1a). Based on what we know about the biology of photosynthesizing cells (in plants, algae, and cyanobacteria), we must predict that transcripts of algal origin, which are supposed to maintain kleptoplast integrity in the slugs, are abundant. Yet, among all RNA sequencing data available for several species (Rumpho et al. 2011; Wägele et al. 2011; Pierce et al. 2012; de Vries et al. 2015), sequencing reads pointing toward transcripts from genes of LGT origin remain close to zero. In fact, they remain well below the counts representing obvious contamination (fig. 2a and b).

Single reads can easily be artifacts. In RNAseq analysis, it is common practice to filter for only those genes that are supported by a reliable number of independent reads. In the RNA-seq analysis by Pierce et al. (2012)—the last RNA-seq report published to claim expression of genes of LGT origin is relevant for kleptoplast performance—the highest read count for a single algal nuclear gene of potential LGT origin was two. Two among 98,238,204 reads. A favored argument to explain why only such few reads are detected is that “the symbiotic chloroplasts resides in only a few cells within the slugs” (Pierce et al. 2012) or that “only a relatively few cells in the slug contain plastids” (Pierce et al. 2015).

Both the appearance of the animal (fig. 1a and b) and factual numbers tell a different story. The chlorophyll content in a 6 mm long *Elysia comogera* is around 3.1 nmol/mg dry weight and in a 10-day-old tomato seedling with fully developed green cotyledons it is about 22.4 nmol/mg dry weight (fig. 2c). A single chloroplast contains 2.5 x 10⁹ chlorophyll (Chl) molecules (Stolz and Walz 1988). One nanomole of Chl thus corresponds to about 2 million chloroplasts, and hence about 6 million chloroplasts are found per mg dry weight in *E. comigera*. Three nanomoles of Chl further translates into a mass of about 2.7 μg, and therefore accounts for about 3% of the total dry weight of slugs. Impressively, even if the estimations would be lower by a factor of 10. Also, if only a few cells of a photosynthetic sacoglossan slug would harbor kleptoplasts, then how would that match up with the concept that photosynthesis continues almost unabated for up to a year to support animal growth (Pierce et al. 2012)? Furthermore, among the RNA-seq data of 2012 from *E. chlorotica*, 4,234 reads for the plastid-encoded *psbA* were detected (Pierce et al. 2012). That is noteworthy, because the samples sequenced were enriched for poly(A)-tailed mRNA prior to sequencing and the plastid mRNA was copurified only as a contamination due to the high AT-content of the plastid transcripts. There is probably even more mRNA encoding *psbA* present than sequenced and even if not, the number of reads for this single *psbA* gene by far exceeds the total number of reads (111) found for the 52 genes of suggested LGT origin. Furthermore, all the reads interpreted to be of LGT origin are ≥99% identical in sequence to the algal transcripts and that would mean they are exempt from evolutionary codon adaptation in the slug’s nucleus.

Slugs analyzed are mostly collected from the wild and then grown on their food alga in open aquaria in the lab. The cultures are not axenic; they cannot be and they do not have to be for the kind of experiments that are currently performed. Contamination of the isolated RNA is unavoidable, but not a problem as long it is monitored. In the most recent transcriptomic analysis on two slug species (de Vries et al. 2015), the number of reads obtained for genes of heterotrophic protists was much higher than those for any algal nuclear gene and they were hence omitted from downstream analysis. We predict the amount of contamination in the data set of *E. chlorotica* (Pierce et al. 2012) to be comparable. The entire sequence data have never been made publicly available, rather only that of the few dozen genes discussed and therefore it was not possible to assess this issue in *E. chlorotica*. But a thought experiment is possible: we 1) do not screen the slug RNA-seq data for potential contamination and we 2) accept the presence of a few mRNA reads as evidence for functional LGT that support kleptoplast maintenance. We could then conclude that hundreds of ciliate genes support kleptoplasty in *E. timida*, but fail to do the same for the short-retaining *E. comigera* albeit present (fig. 2a). The most rational conclusion that remains is that RNA-seq offers no support for the expression of slug nuclear genes that originate from the food alga. These slugs are not what they eat, and they eat a lot (Christa et al. 2014).

Recently, evidence for algal LGT in Sacoglossa other than sequencing data emerged: a study by Schwartz et al. (2014) used fluorescence in situ hybridization (FISH) to localize genes of algal origin among slug chromosome spreads. That report was quickly picked up by the popular press and it currently scores among the top 5% of all articles so far evaluated by Altmetric.com. Evidently, the public cares a lot about LGT in slugs. The public, however, is likely less aware that FISH analysis on two slug species (de Vries et al. 2015), sequencing reads pointing toward transcripts from genes of LGT origin remain close to zero. In fact, they remain well below the counts representing obvious contamination (fig. 2a and b).

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from these technical issues, FISH analysis is not a suitable tool for providing evidence for LGT. The only reliable evidence for LGT would be to demonstrate the integration of algal DNA into the context of slug chromosomes (through DNA sequencing), from where it is expressed to support the stolen organelles by the product being specifically targeted to kleptoplasts.

And although independent genome data of *E. chlorotica* is available (Pierce et al. 2012) to the authors of the FISH analysis, it has not been used to support their concept and also challenges published slug genome data that found no evidence for algal LGTs in *E. chlorotica* (Bhattacharya et al. 2013).

**Fig. 1.**—Sacoglossan slugs can house millions of kleptoplasts. (a) Shown are two of the earliest depictions of sacoglossan slugs and their “green” digestive tubules that can pervade the entire body. On the left a drawing by van Hasselt from 1824 showing *Plakobranchus ocellatus* and on the right a drawing of *Elysia viridis* by J. Thomas from 1852. Both demonstrate that an extensive digestive system, able to house millions of kleptoplasts, is not limited to *Elysia chlorotica*. Note how the digestive tubules of *E. viridis* pervade even the head of the animal. (b) The extent of stored plastids becomes apparent when viewing the slugs (here *Elysia timida*) under the microscope and filtering for the chlorophyll autofluorescence of the kleptoplasts (red-orange hue). In the middle, a detail of a region of the parapodia, with the individual digestive tubules being visible through the kleptoplasts’ fluorescence. Zooming in further reveals the density with which the kleptoplasts are packed into the cytosol of the cells forming the digestive tubules.
Algal Genes and Photosynthetic Slugs

As a last word on LGT, it should be mentioned that LGT to euukaryotes is manifest in two fundamentally different forms. First, there is gene transfer from organelles to the nucleus, or also called EGT. EGT is a continuous, ongoing process, and incontrovertibly documented in all sequenced genomes of euukaryotes (Timmis et al. 2004; Hakkan-Covo et al. 2010; Boto 2014). Second is outright LGT, where the donors are not chloroplasts or mitochondria. Newer findings show that latter, though it does occasionally occur, is extremely rare and does not manifest itself in the bigger picture of euukaryotic evolution (Ku et al. 2015). In this context, it is important to note that long-term retention of kleptoplasts evolved several times independently in sacoglossan sea slugs (Maeda et al. 2010; de Vries, Christa, et al. 2014; Christa et al. 2015). Hence, if the expression of nuclear genes of algal origin is the reason for robust kleptoplasts in one species, then the same should apply to other species as well. If all Sacoglossa retained functional kleptoplasts in a LGT-dependent manner, then they would have to be the record holders for LGT among animals, their LGT events outnumbering all other cases in animals thus far reported. Are Sacoglossa LGT magnets? Neither genome nor transcriptome data from these animals indicate that to be the case. Occam’s razor dictates favoring a less assumptive scenario.

Stable Kleptoplasts in the Absence of LGT

Since the 1970s, it is known that some plastids sequestered by the sea slugs show a remarkable independent robustness (Giles and Sarafis 1972; Trench and Ohlhorst 1976; Green et al. 2005). The best explanations we have for robust plastids are effective photoprotection mechanisms (Serôdio et al. 2014; Cruz et al. 2015), a different coding capacity of the plastid genomes in question (Rumpho et al. 2000; de Vries et al. 2013) and maybe an overall difference in the stability (half-life) of essential proteins. That plastids sequestered by the slugs are intrinsically robust is, based on current information, the most parsimonious scenario. It explains how such a broad range of slug species can perform kleptoplasty (Christa et al. 2015) and why plastids of the same source can behave identically in slug species that differ in their ability to survive food deprivation (de Vries et al. 2015). Slugs acquiring robust plastids will not automatically retain them long-term and endure starvation as recently interpreted (Pierce et al. 2015). It is of equal importance that the slugs are physiologically adapted and require to retain them functionally (de Vries, Rauch, et al. 2014). This likely depends on whether they experience food deprivation (de Vries et al. 2015). Slugs acquiring robust plastids will not automatically retain them long-term and endure starvation as recently interpreted (Pierce et al. 2015). It is of equal importance that the slugs are physiologically adapted and require to retain them functionally (de Vries, Rauch, et al. 2014). This likely depends on whether they experience food deprivation in their habitat due to seasonal variation or not (Cruz et al. 2013; Wägele and Martin 2013; de Vries et al. 2015).

The ability to sequester and maintain an entire heterologous structure of foreign origin is not restricted to sacoglossa and their plastids. For the purpose of using them as a

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Figure 2: Among RNA-seq data, contaminating reads exceed reads of algal origin. (a) At the top the total number of reads (in million, M) recently sequenced for *Elysia cornigera* and *Elysia timida* (de Vries et al. 2015) are shown. Those reads were assembled into contigs and all contigs subjected to a BLASTx-based distribution analysis against RefSeq to determine their distribution among the taxonomic groups listed. Note that 1) the number of reads of protist origin in all cases exceeds those of green algal origin and 2) in the LtR species of reads of protist origin in all cases exceeds those of green algal origin and 3) the number of reads of protist origin in all cases exceeds those of green algal origin. (b) Among RNA-seq data, contaminating reads exceed reads of algal origin. (c) Green leaves and Green Sacoglossa.
defensive organ, some aelolidioidean sea slugs incorporate cni-
docysts from their cnidarian prey to expose them on their
surface (Obermann et al. 2012). Similar to the kleptoplasts,
cnidocysts are first incorporated through oral feeding and as
part of the regular diet. The peculiar thing is the release of the
kleptoplasts from the phagosomes into the cytosol of the di-
gestive epithelial cells. The latter appears more common when
only organelles and not entire symbiotic organisms are re-
tained by a host. The ciliate Myrionectra rubra releases tran-
scriptionally active nuclei and plastids of its prey algae into the
cytosol (Johnson et al. 2007), whereas symbiotic Chlorella
algae of Paramecium bursaria or Hydra viridis remain inside
a specialized digestive vacuole and isolated from the host’s
cytosol (Nowack and Melkonian 2010; Fujishima and
Kodama 2012). It is not known how the plastids are specific-
ally sorted from other food particles and then released into
the cytosol or really why. Does it facilitate the easier ex-
change of substrate and metabolites? These observations, to-
gether with how Sacoglossa deal with kleptoplasm-produced toxins
such as reactive oxygen species and the general differences in
stavation tolerance, remain promising research topics. All
of these, however, are not associated with LGTs of algal
origin.

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