

Review

From famine to feast? Selecting nuclear DNA sequence loci for plant species-level phylogeny reconstruction

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Phylogenetic analyses of DNA sequences have prompted spectacular progress in assembling the Tree of Life. However, progress in constructing phylogenies among closely related species, at least for plants, has been less encouraging. We show that for plants, the rapid accumulation of DNA characters at higher taxonomic levels has not been matched by conventional sequence loci at the species level, leaving a lack of well-resolved gene trees that is hindering investigations of many fundamental questions in plant evolutionary biology. The most popular approach to address this problem has been to use low-copy nuclear genes as a source of DNA sequence data. However, this has had limited success because levels of variation among nuclear intron sequences across groups of closely related species are extremely variable and generally lower than conventionally used loci, and because no universally useful low-copy nuclear DNA sequence loci have been developed. This suggests that solutions will, for the most part, be lineage-specific, prompting a move away from ‘universal’ gene thinking for species-level phylogenetics. The benefits and limitations of alternative approaches to locate more variable nuclear loci are discussed and the potential of anonymous non-genic nuclear loci is highlighted. Given the virtually unlimited number of loci that can be generated using these new approaches, it is clear that effective screening will be critical for efficient selection of the most informative loci. Strategies for screening are outlined.

Keywords: Tree of Life; plant phylogeny; nuclear DNA sequence loci; low-copy nuclear gene; comparative anchor tagged sequence; sequence characterized amplified region

1. INTRODUCTION

Reconstructing the Tree of Life has been a central objective of evolutionary biology since phylogenetic trees were first proposed as a way of representing evolutionary relationships. Progress towards that goal has accelerated in the last two decades and assembling the Tree of Life is now the focus of more research than at any time over the past 150 years (e.g. Soltis & Soltis 2001; Donoghue & Cracraft 2004; Palmer *et al.* 2004). This has been prompted by theoretical and methodological advances, vastly increased computational power, and technical improvements in DNA sequencing that enable the generation of large volumes of character data relatively quickly and cheaply. It is now possible to build robust hypotheses of relationships for large numbers of taxa and to contemplate the vision of a complete Tree of Known Life (Soltis & Soltis 2001; Watanabe 2002; Donoghue & Cracraft 2004). Current interest in phylogeny reconstruction has also been driven by ever wider application of phylogenetic trees within as well as beyond the confines of systematics and evolutionary biology (e.g. Soltis *et al.* 1999; Savolainen & Chase

2003; Futuyma 2004; Yates *et al.* 2004). Phylogenetic trees provide not only the basis for classification, but also studies of character evolution (Schultheis & Baldwin 1999), hybridization (e.g. Hughes *et al.* 2002; Linder & Rieseberg 2004), polyploidy (e.g. Doyle *et al.* 2003b, 2004), biogeography (e.g. Lavin *et al.* 2004; Pennington *et al.* 2004), origins of domestication (e.g. Wang *et al.* 1999; Nesbitt & Tanksley 2002) and speciation and species diversification (e.g. Barraclough & Vogler 2000; Barraclough & Nee 2001). Phylogenies also provide information to develop comprehensive comparative systems in developmental biology (e.g. Doust & Kellogg 2002; Thießen *et al.* 2002) and comparative genomics (e.g. Bennetzen & Kellogg 1997; Eisen & Fraser 2003; Hong *et al.* 2003). In the last two decades, phylogenies have come of age as a universal component of comparative biology (Hillis 2004).

Questions remain about the best ways to measure progress towards estimating the Tree of Life (Donoghue 2004). Numbers of published phylogenetic trees and taxa represented therein alongside measures of levels of resolution among taxa, statistical confidence in published trees (Sanderson 1995), and discovery of paraphyly, polyphyly, and circumscription of monophyletic groups, all suggest that recent progress has

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indeed been impressive (reviewed in Cracraft & Donoghue 2004). However, measures of success are greatly influenced by the intended use(s) of the resultant phylogenetic trees. For the purposes of ridding classifications of non-monophyletic groups, completely resolved trees with uniformly high statistical support for all nodes are not necessarily required. Analyses of exemplar taxa that resolve a subset of well-supported nodes will often suffice for the purposes of providing robust phylogenetic classifications. There is little doubt that in this sense there has been massive progress towards assembling the Tree of Life. The angiosperms provide a good example. Despite some persistent areas of poor resolution and/or weak support (e.g. Wortley *et al.* 2005), the branching order of most clades of angiosperms is now relatively clear and well supported (e.g. Soltis *et al.* 1999; Savolainen & Chase 2003; Soltis & Soltis 2004), prompting a new classification (APGII 2003). Similar progress has been made towards delimiting genera and understanding relationships within many angiosperm families (e.g. Lavin *et al.* 2001). In this realm there are clear theoretical reasons to believe that further progress can be made by simply increasing the number of characters via sequencing additional readily accessible loci that can be combined in simultaneous analyses to increase resolution, accuracy (Hillis 1996, 1998) and support (Bremer *et al.* 1999) for critical nodes (Wortley *et al.* 2005). This is borne out by a number of empirical studies that have employed large scale DNA sequence datasets (Herniou *et al.* 2001; Baptiste *et al.* 2002; Matsuoka *et al.* 2002; Rokas *et al.* 2003), albeit with the provisos that increased character sampling is not compromised by sparser taxon sampling (Soltis *et al.* 2004) and that the methodological challenges posed by increasingly large data matrices can be adequately addressed (Sanderson & Driskell 2003).

However, when we come to examine progress in constructing accurate phylogenies among closely related species the situation, at least for plants, is less encouraging. Partially resolved gene and species trees are of limited use for studies of hybridization and polyploidy, character evolution, species diversification, speciation and domestication, or as comprehensive comparative systems underpinning biology more generally. Species-level phylogenies for detailed evolutionary studies often demand not just complete or near-complete taxon sampling, but ideally multiple accessions within species, as well as complete or near-complete resolution and high statistical support. Accurate statements about hybrid and polyploid origins require that the terminal branches leading to putative hybrids and their parents on gene trees are resolved and well supported. For example, lack of resolution in gene trees derived from conventional cpDNA and nrDNA loci in the legume genus *Leucaena* (figure 1) meant that the parentage of hybrids and polyploid species in the genus could not be hypothesized beyond major clades using these loci (Hughes *et al.* 2002). The same applies to studies of domestication where gene trees are used to infer potential progenitors of crop plants (e.g. Emshwiller & Doyle 1999, 2002). Using phylogenies to understand geographical patterns of species diversification or to

plot lineage diversification through time and understand speciation processes requires similarly high-quality phylogenetic trees (e.g. Barraclough & Nee 2001).

Conflict attributable to lineage sorting, hybridization, polyploidy and introgression can cause gene genealogies to differ from the branching history of the organisms from which the genes were sampled (Pamilo & Nei 1988; Doyle 1992; Hillis 1995; Wendel & Doyle 1998). The impacts of these processes are thought to be more serious at species level than at higher levels, suggesting that fully resolved divergent species trees should not necessarily be expected for many plant genera (e.g. Linder & Rieseberg 2004). This means that the general desirability of employing multiple independent loci to infer accurate phylogenies can be much more critical in species-level phylogenies especially for plant groups where reticulation is common (Ferguson & Sang 2001; Raymond *et al.* 2002; Linder & Rieseberg 2004; Hegarty & Hiscock 2005). Employing additional DNA sequence loci to obtain greater species-tree resolution works on the premise that individual matrices will either be combined for simultaneous analysis to maximize congruence among independent sources of data (Nixon & Carpenter 1996), or, in the case of reticulation due to hybridization-introgression, that sets of congruent genes can be used to distinguish divergent from reticulate relationships. Differentiating between these alternatives requires gene trees with sufficient resolution to discriminate conflict from congruence.

Here we attempt to encapsulate how far plant species-level phylogenetics has progressed. We show that the rapid accumulation of DNA characters at higher taxonomic levels has not typically been matched in species-level analyses. It is also apparent that success in locating DNA sequence loci that can resolve relationships among species has been extremely patchy, lagging behind studies of animal taxa. Examples of densely sampled, well-resolved and supported species-level phylogenies for plants based on multiple independent loci are scarce and in many cases extremely challenging to reconstruct (see below). This means that many interesting questions in plant evolutionary biology and biogeography are currently frustrated by lack of resolution towards the tips of the tree. There has been little mention of these issues in the recent discussions about completing the Tree of Life (Cracraft & Donoghue 2004), although there have been recurrent calls for improved species-level phylogenies from those involved in evolutionary studies (e.g. Baldwin & Sanderson 1998; Barraclough & Nee 2001; Doyle *et al.* 2003b; Bailey *et al.* 2004; Futuyama 2004; Linder & Rieseberg 2004). In the last few years, new approaches to identify nuclear DNA sequence loci have emerged that have the potential to transform the current famine into a veritable feast of hypervariable species-level loci. We provide an explanatory overview that describes and compares the utility, potential benefits and pitfalls of the different approaches. Finally, we discuss the need to implement pilot studies that involve the screening of exemplar accessions to rank loci by potential utility prior to embarking on full scale sequence analysis of any one locus.

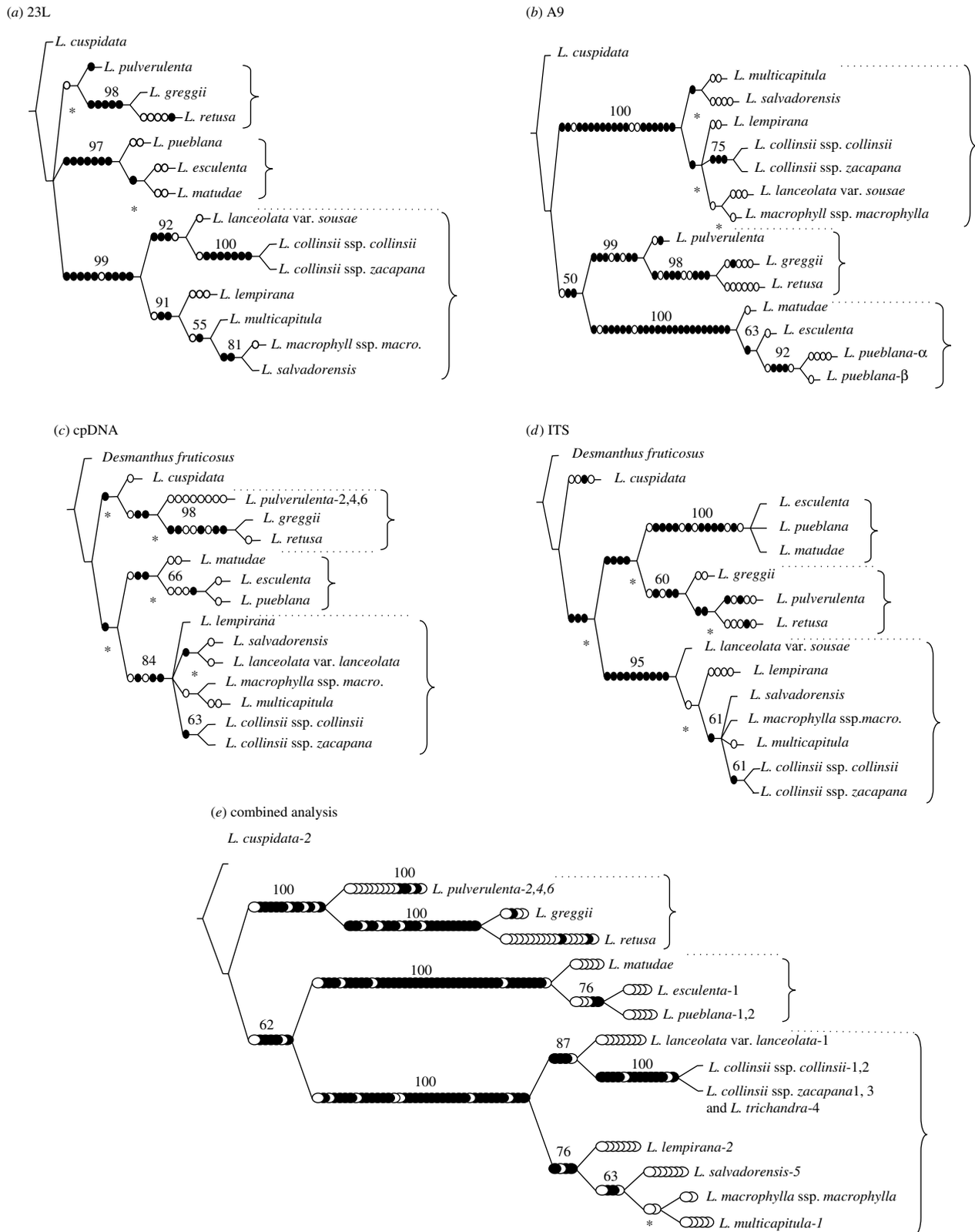


Figure 1. Example of increased resolution and support obtained among closely related diploid species of the Mimosoid legume genus *Leucaena* using a SCAR-based approach (modified and extended from Bailey *et al.* 2004). A fully resolved and well-supported diploid species tree for *Leucaena* is required to investigate the origins of a set of putative hybrid and five polyploid species (Hughes *et al.* 2002). Solid and open circles along branches represent unique and homoplastic unambiguous character state transformations, respectively. Values above branches are strict consensus bootstrap support values; nodes that are not present in the strict consensus trees are marked with an *. (a) Anonymous SCAR-based nuclear locus 23L—one of four EMPTs; $L=71$; CFI=0.75; CI=0.80; RI=0.88. (b) Anonymous SCAR-based nuclear locus A9—one of six EMPTs; $L=132$; CFI=0.69; CI=0.75; RI=0.88. (c) Chloroplast RFLP—one of 24 EMPTs; $L=63$; CFI=0.46; CI=0.58; RI=0.69. (d) nrDNA ITS—one of six EMPTs; $L=100$; CFI=0.46; CI=0.70; RI=0.81. (e) Combined analysis of 23L, A9, cpDNA and ITS datasets—one of three EMPTs; $L=339$; CFI=0.92; CI=0.73; RI=0.84. EMPT = equally most parsimonious trees; L = length (number of steps); CFI = consensus fork index (calculated from strict consensus trees); CI = consistency index; RI = retention index.

2. SURVEY OF SPECIES-LEVEL MATRICES

One of the strengths of DNA sequence data for resolving relationships is the scope to select among conserved and more rapidly evolving sequence regions, the 'tortoise and hare' of Small *et al.* (1998), to address questions at different hierarchical levels (e.g. Small *et al.* 1998, 2004; Soltis & Soltis 1998; Yang 1998). Molecular systematists have thus been able to tailor the selection of sequence regions to the question at hand—using more slowly evolving loci to analyse higher level relationships and more rapidly evolving regions for studies of closely related species. In this way, sequences from different genomes, genes and coding/non-coding regions can potentially generate data appropriate to resolve all branches of the Tree of Life from species level (or even within species) upwards.

There are a number of different ways to measure the potential informativeness of different DNA sequence loci (Wortley & Scotland *in press*). Two measures, the number of parsimony informative (PI) characters and the percentage of PI characters (total number of PI characters divided by the aligned length), are widely used. Wortley & Scotland (*in press*) point to the minimum number of PI character state changes as a more accurate measure of potential utility. However, this measure is rarely reported and for comparisons among molecular datasets is strongly correlated with number of PI characters. Number and percentage PI characters are not independent. From a tree building perspective, the total number of PI characters is more important than the per cent variability, in that a highly variable but very short region may not provide sufficient characters. However, from a practical perspective, very long regions, while potentially contributing more characters, may be inefficient in terms of sequencing effort. In spite of falling costs of sequencing, obtaining the maximum number of characters per sequencing reaction will remain an important factor for species-level phylogeny reconstruction given the scale of taxon sampling with multiple accessions of species that is needed. Thus, both total number of PI characters and per cent PI characters are useful measures for comparing loci. These can be reported relative to the number of taxa as character–taxon ratios.

We have undertaken a two-tier survey of recent plant species-level phylogenetic analyses. First, we surveyed a sample of the most recently published studies (2003 and 2004) from *American Journal of Botany*, *Molecular Phylogenetics and Evolution*, *Systematic Botany* and *Taxon* (136 studies encompassing 345 individual data matrices). These were assessed in terms of their objectives, taxon sampling, character sampling (numbers and types of DNA sequence loci used) and the degree of resolution obtained. In order to assess resolution we used the consensus fork index (CFI; Colless 1980; the number of resolved nodes on a strict consensus tree divided by the number of possible resolved nodes, $n-2$, where n is the number of terminals). Second, we looked in more detail at potential informativeness of different types of sequence loci using a subset of 226 recent plant species-level data matrices from studies that include nuclear (nDNA) and non-coding chloroplast (cpDNA) and/or nuclear ribosomal (nrDNA) loci. We have restricted both surveys

and discussions to sets of congeneric species, and data matrix statistics to ingroup species only. This could be viewed as somewhat arbitrary not least because genera are not uniformly applied, but also because some studies focus on subclades within genera. Numbers of PI characters can also be influenced by levels of taxon sampling within genera. Nevertheless, a broad sample of species-level studies allows for the direct comparison of variation provided by different classes of loci to evaluate the general utility of each class.

3. CHLOROPLAST AND ITS DATA

Results from the broad survey of species-level matrices are summarized in figure 2. So far, the vast majority of species-level phylogenetic analyses of plants have relied on a limited set of non-coding cpDNA loci and the nrDNA internal transcribed spacer (ITS) region (ITS 1, 5.8S and ITS 2; figure 2a; Alvarez & Wendel 2003; Shaw *et al.* 2005b). These loci account for 87% of 345 matrices surveyed (figure 2a) and have been favoured because they are variable at low taxonomic levels and easy to amplify using universal primers (Taberlet *et al.* 1991; Baldwin *et al.* 1995; Shaw *et al.* 2005b). Routine application of these loci has produced an explosion of new phylogenetic data providing insights into species relationships across a wide range of plant genera. The average resolution across studies as measured by the CFI is 0.64 (figure 2c). Even at this level of resolution, many of these analyses serve the primary purpose—i.e. form a basis for generic delimitation and infrageneric classification (figure 2b; Alvarez & Wendel 2003). However, the limitations of relying on this limited set of loci are also apparent, especially for studies with objectives beyond classification (figure 2b). Only 11% (15 trees) of our sample of recently published species-level phylogenies are fully resolved, more than half are poorly resolved (CFI < 0.6), and 14% are very poorly resolved (CFI < 0.4; figure 2c). Lack of resolution is, thus, a widespread problem even for morphologically diverse species representing large genera (Wojciechowski *et al.* 1999; Richardson *et al.* 2001; Malcomber 2002; Mitchell & Heenan 2002; Syring *et al.* *in press*). Usually, this lack of resolution is directly attributable to insufficient variation (Small *et al.* 1998; Bailey *et al.* 2004; Shaw *et al.* 2005b), rather than incongruence, although poorly resolved trees can in themselves preclude observation of incongruence. It is also worth noting that taxon sampling is extremely variable across species-level studies, with only 37% fully sampled (figure 2e).

Results from the more detailed comparisons of the potential utility of different DNA sequence loci are presented in figure 3. Percentage PI character values for non-coding cpDNA range from 0.2 to 8.5 (–13.1 in Shaw *et al.* 2005b) and for ITS from 0.3 to 14 (–24) (figure 3). These values are an order of magnitude lower than those available in many higher level analyses (typically 20–60%). They are also lower than commonly used mitochondrial (mt) DNA loci in animals, where highly variable (typically 15–30% PI characters) DNA sequences have facilitated reconstruction of highly resolved species-level phylogenies (Avise *et al.* 1987; Moritz *et al.* 1987; Moore 1995) and speciation events (e.g. Irwin *et al.* 2001). The low percentages of

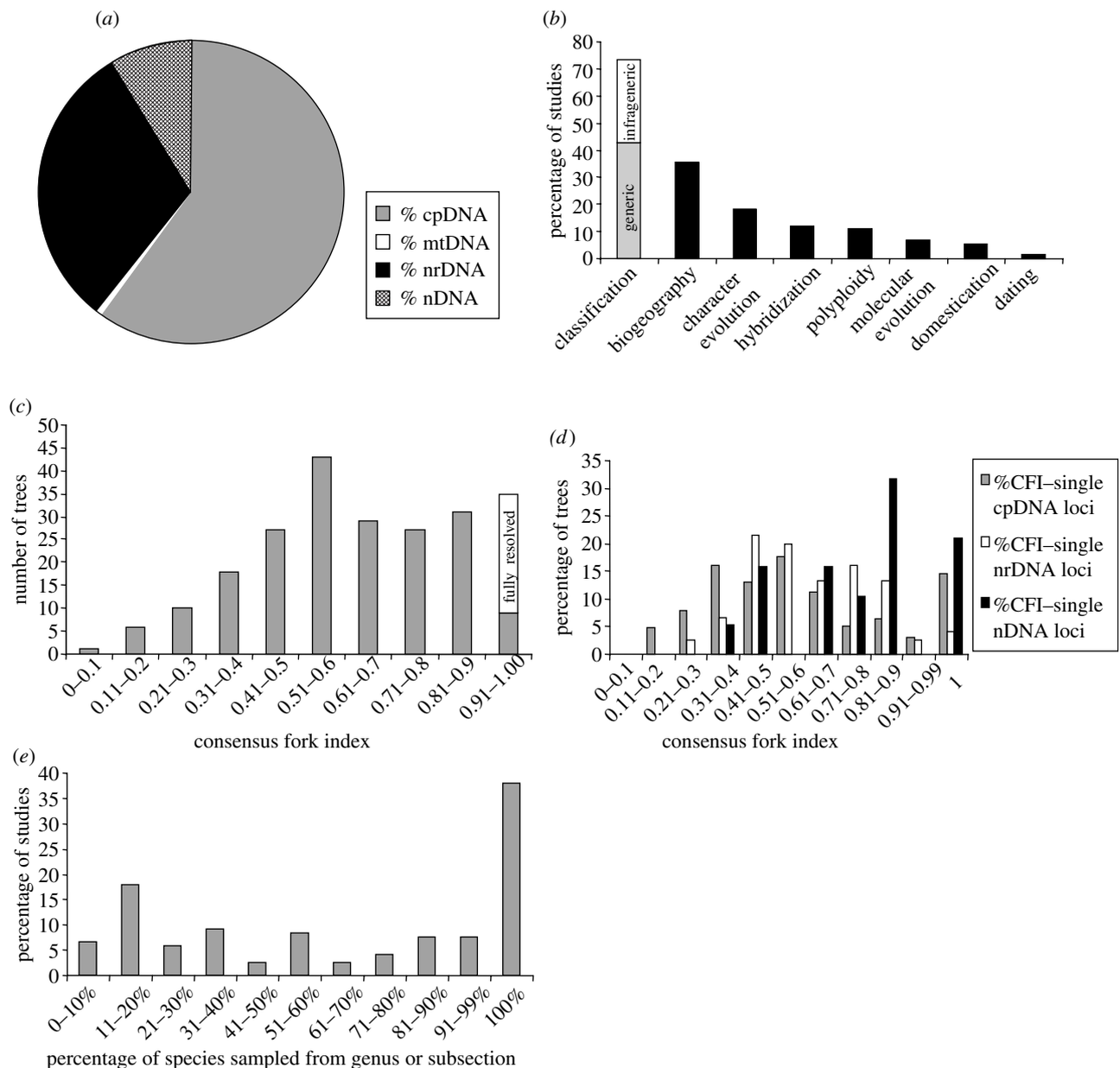


Figure 2. Attributes of current plant species-level phylogenetic studies based on a survey of 345 data matrices from 136 studies (see text for details). (a) Loci currently used in species-level analyses: proportions of cpDNA, nrDNA (ITS), mtDNA and nDNA loci. (b) Objectives and applications of species-level phylogenetic analyses. (c) Resolution obtained in recent species-level analyses across all loci: frequency distribution of consensus fork indices, including both combined and separate analyses. (d) Resolution obtained using different loci: frequency distribution of consensus fork indices for analyses of individual loci. (e) Variation in taxon sampling in current species-level analyses.

PI characters are reflected in lower CFI values for cpDNA and ITS loci when analysed alone compared to some nuclear loci (figure 2d).

In the face of insufficient variation to resolve relationships among sequences from conventional non-coding cpDNA and/or ITS loci, a number of approaches have been adopted (see below). Increasing the amount of cpDNA sequence data, potentially guided by selecting more variable non-coding cpDNA loci (Shaw *et al.* 2005b), has been successfully used to obtain greater resolution and support. Inevitably, this approach involves large volumes of DNA sequencing (e.g. 4–7 kb of plastid sequence used by Cronn *et al.* 2002; Clarkson *et al.* 2004; Shaw & Small 2004) and still does not guarantee well-resolved trees (e.g. Shaw & Small 2004). Other approaches have

been to sequence the structurally complex external transcribed spacer (ETS; e.g. Linder *et al.* 2000), or resort to dominant PCR-based fragment length characters derived from amplified fragment length polymorphisms (AFLPs), inter-simple sequence repeats (ISSRs) or randomly amplified polymorphic DNAs (RAPDs) (reviewed by Wolfe & Liston 1998; Harris 1999). Contrary to the situation in the animal kingdom, mtDNA sequences have only played a minor role in phylogenetic studies of plants. This is because of structural instability, gene transfer to the nucleus and alleged sequence conservation (Wolfe *et al.* 1987; Palmer 1992; Soltis & Soltis 1998). However, some recent studies have found mtDNA sequence loci sufficient to build species-level phylogenies in plants (e.g. Sanjur *et al.* 2002).

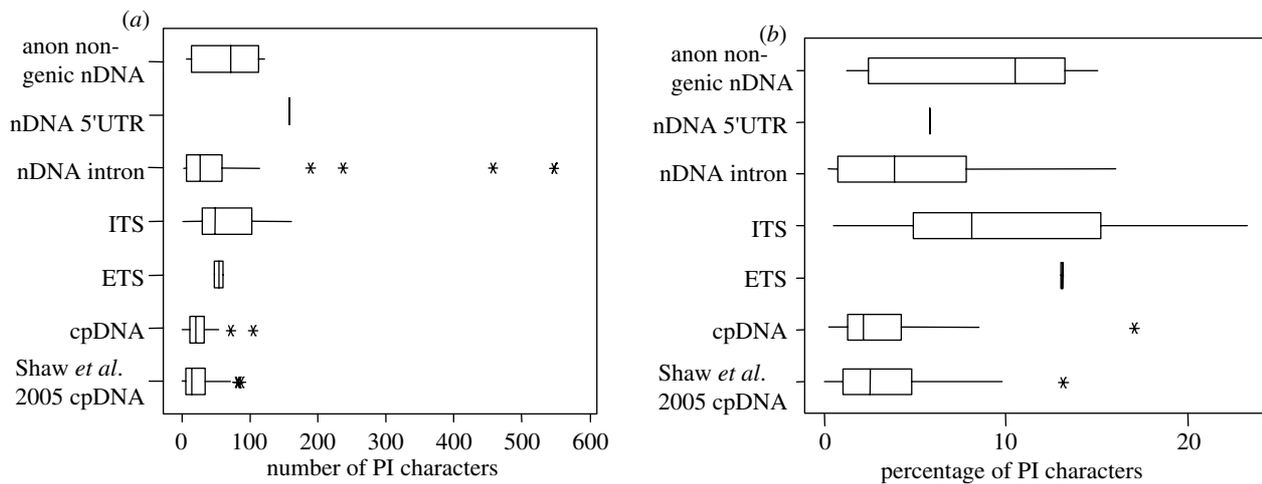


Figure 3. Potential informativeness of cpDNA, nrDNA ITS, nrDNA ETS, nDNA intron, nDNA 5' UTR and anonymous, SCAR-based DNA sequence loci at species level. (a) Number of PI characters per locus. (b) Percentage of PI characters for individual loci. Boxes encompass the 75% interquartiles, showing the median while outliers are indicated with an *. Data are derived from species-level phylogenetic analyses of 226 matrices from studies of plant genera. Character statistics are calculated for congeneric terminals only, excluding outgroup taxa. Tallies of characters included both PI substitutions and PI indels coded according to the simple gap coding method of Simmons & Ochoterena (2000). Plant groups included are: *Sphagnum* (Shaw *et al.* 2003), *Leucaena* (Bailey *et al.* 2004 and unpublished data), *Gossypium* (Cronn *et al.* 2002), *Gaetnera* (Malcomber 2002), *Spaerocardamum* (Bailey & Doyle 1999), *Passiflora* (Yockteng & Nadot 2004), *Glycine*—alignments assembled for nine species based on data from Doyle *et al.* (1996, 2003a,b) and Sakai *et al.* (2003), *Paeonia*—alignments assembled for eight species based on data from Sang *et al.* (1995, 1997), Ferguson & Sang (2001) and Tank & Sang (2001), *Lycopersicum*—alignments assembled for 13 accessions of seven species based on data from Nesbitt & Tanksley (2002), *Viburnum* (Winkworth & Donoghue 2004), *Clarkia* (Ford & Gottlieb 2003), *Hibiscus* (Small 2004), *Neillia/Stephandra* (Oh & Potter 2003) and *Bursera* (Weeks & Simpson 2004). The broader survey of non-coding cpDNA loci (Shaw *et al.* 2005b) is included as a second cpDNA bar.

Aside from the lack of sufficient variation in many groups, cpDNA and nrDNA loci individually are of reduced utility for reconstructing relationships among potentially reticulating species (Linder & Rieseberg 2004). Uniparental inheritance of plastids makes cpDNA essential for teasing apart maternal versus paternal contributions to putative hybrid species when compared to other data. Similarly, nrDNA multi-copy sequences that have undergone complete concerted evolution do not retain evidence of biparental inheritance and therefore reticulation (Baldwin *et al.* 1995; Wendel *et al.* 1995). The apparently unpredictable extent and direction of concerted evolution among nrDNA repeats, which can even vary among individuals of the same species (Doyle *et al.* 2004), can create situations where nrDNA data provide partial and potentially confusing information about reticulation. Conversely, lack of concerted evolution can create serious paralogy-related problems in some ITS datasets (Alvarez & Wendel 2003; Bailey *et al.* 2003; Doyle *et al.* 2004). These limitations mean that multiple independent bi-parentally inherited loci are essential for species-level phylogenetic studies where reticulation is a possibility (Ferguson & Sang 2001; Raymond *et al.* 2002; Linder & Rieseberg 2004; Hegarty & Hiscock 2005), and these can only come from the nuclear genome.

4. STRATEGIES FOR SELECTING NUCLEAR SEQUENCE LOCI

(a) *Low-copy nuclear gene approaches*

The most widely used approach to resolve relationships among closely related plant species where cpDNA and ITS loci fail to provide resolution has been to use

sequence data from low-copy nuclear genes (LCNG), or specific members of multi-gene families and especially their introns (Doyle & Doyle 1999; Cronn *et al.* 2002; Sang 2002; Small *et al.* 2004). Initial studies using nuclear sequences were restricted by the availability of primers to a handful of LCNGs (e.g. floral development genes) under investigation in related taxa in other fields (Strand *et al.* 1997; Small *et al.* 1998; Cronn *et al.* 2002). This approach has succeeded in locating sporadic highly variable loci for specific plant groups (e.g. Histone H3-D in *Glycine*, Doyle *et al.* 1996; GBSSI/*waxy* in grasses, Mason-Gamer *et al.* 1998 and Rosaceae, Evans *et al.* 2000; GPAT in *Paeonia*, Tank & Sang 2001; *pgiC1* and *pgiC2* in *Clarkia*, Ford & Gottlieb 2003; *cycloidea* in *Lupinus*, Ree *et al.* 2004). However, LCNG loci still comprise less than 10% of published species-level matrices (figure 2a), and for the vast majority of plant genera choice of LCNG loci remains limited and access to alternatives a significant hurdle.

Also, it is increasingly clear that nuclear-encoded loci vary widely in nucleotide substitution rates among closely related species. For example, Cronn *et al.* (2002) found a fivefold range in substitution rates among 12 nuclear encoded loci developed for *Gossypium*. In combined analysis of these nuclear loci plus ITS, 53% of the PI characters were recovered from just two loci (ITS and Fad2-1). Furthermore, the ITS region provided 33% of PI sites from just 10% of the total sequence data. An even wider sevenfold difference in substitution rates was documented in a survey of 36 nuclear genes for the same species (Senchina *et al.* 2003). There are clear indications of similar variability in rates of evolution among intron sequences of closely

related species from other genera (e.g. Doyle *et al.* 2003a, 2004 for *Glycine*; Syring *et al.* in press for *Pinus*). A wider survey of nuclear intron datasets confirms this variability which ranges from 0.21 to 16% PI characters across groups of congeneric species (figure 3). Of the 24 nuclear intron datasets surveyed here which have ITS data for the same taxa, only seven provided more PI characters than ITS (figure 3). While this variability has been recognized for some time (Small *et al.* 2004), what has not been widely acknowledged is that the majority of nuclear introns sequenced across groups of species, so far, are less variable (and often much less variable) and provide fewer PI characters than ITS. Only a small subset exhibits similar or higher levels of variation (figure 3). This is perhaps not surprising considering that some introns, and particularly large introns, have been shown to influence patterns of gene expression, acting as *cis*-regulatory elements (e.g. Sieburth & Meyerowitz 1997). A similarly wide span of variation has been found for LCNG loci used in phylogenetic analyses of animal species where most LCNG sequences are less variable than mtDNA loci (e.g. Helbig *et al.* 2005; Peters *et al.* 2005).

It is also notable that no universally useful LCNG sequence loci have been developed during the last 10 years (Sang 2002). Low-copy nuclear loci have had to be developed specifically for the taxonomic group of interest. In fact, building a phylogeny for a particular plant group has been the primary goal of many of the studies that have also discussed the potential broader utility of specific genes (e.g. Doyle *et al.* 1996; Galloway *et al.* 1998; Mason-Gamer *et al.* 1998; Small *et al.* 1998; Bailey & Doyle 1999; Emshwiller & Doyle 1999; Ree *et al.* 2004), a potential that has not so far been realized. Difficulties of extrapolating loci that have provided informative variation in one group can be due to failure of primers to amplify, variation in copy number, differences in intron presence or length, or differences in nucleotide substitution rates in different plant groups. Thus, up to now it seems that development of LCNG sequence loci has been somewhat of a lottery spawning the occasional lucky jackpot against a backdrop of generally disappointing results and considerable investment in developing and optimizing loci that often fail to produce much data (figure 3). This backdrop is rarely reported in the literature but is reflected in the experiences of several researchers who have acknowledged an element of good fortune in developing highly informative LCNG sequence loci, when they come to try to find similarly variable additional loci for the same group of plants (Doyle *et al.* 2003a; Small *et al.* 2004).

A further issue of concern here is the potential imbalance caused by one or a few high variability loci dominating and potentially distorting analyses. In comparisons of closely related species, the range of variation between loci is often skewed towards having few characters per locus. Having one highly variable LCNG locus alongside a set of much less variable loci may create a false sense of security when reconstructing species trees because poorly resolved trees are less likely to reveal incongruence. Furthermore, the combination of such loci in simultaneous analysis runs an increased risk of having one or a few highly variable loci that are

incongruent with the underlying species tree negatively impacting the signal provided by less variable loci (Bull *et al.* 1993; Miyamoto & Fitch 1995; Page & Charleston 1997; Slowinski & Page 1999; Page 2000). Methods that screen multiple loci for comparable levels of informative variation prior to generating fully sampled trees (see below) will ensure that individual loci provide more equitable contributions towards combined analyses. While it is worth bearing in mind the idea that different loci may provide resolution on different parts of the tree (e.g. Pennington 1996; Ree *et al.* 2004; Helbig *et al.* 2005), this has not been convincingly demonstrated and balanced contributions of character data from a set of highly variable individual loci is likely to be more informative.

(b) Comparative anchor tagged sequence-based approaches

In the absence of universally accessible and informative LCNG loci, alternative approaches to locate more variable nuclear genes have been tried. The first of these involves comparisons of EST and/or complete genome sequences between model organisms to identify evolutionarily conserved regions, termed comparative anchor tagged sequences (CATS) (Lyons *et al.* 1997; Chandappa *et al.* 2005; Syring *et al.* in press) or conserved orthologue set (COS) markers (Fulton *et al.* 2002). While the primary goal of many of these studies has been comparative genome mapping, CATS markers can be used to develop sets of primers circumscribing potentially amplifiable sequence regions for phylogenetic analyses (figure 4a). For example, comparison of the tomato EST database with the *Arabidopsis* genome sequence identified a set of more than 1000 COS markers (Fulton *et al.* 2002). Similarly, screening of the *Medicago trunculata* EST database against the *Arabidopsis* genome sequence, and other available legume sequence, identified 274 loci that show strong sequence similarity. A subset of these are being screened as potential DNA sequence loci for species-level phylogenetic reconstruction of the large legume genus *Astragalus* (Scherson *et al.* in press). Similarly, Chandappa *et al.* (2005) used genome and EST sequence alignments to design sets of conserved markers for monots based on rice–onion and rice–banana alignments. This approach has also been used directly with the *Arabidopsis* and rice genome sequences to identify conserved DNA oligomers that can be employed as primers to amplify orthologous DNA sequence loci for species-level phylogenetics (Padolina *et al.* 2004; Xu *et al.* 2004). One of the most notable features of these methods is the large number of potential primer pairs that can be generated. The few studies undertaken so far identify 100s or 1000s of possible DNA sequence loci and primer pairs. At the extreme, 13 418 candidate primer pair combinations were identified in the studies of Padolina *et al.* (2004) and Xu *et al.* (2004).

Development of CATS-based approaches is at an early stage and, as far as we are aware, they have yielded very few, if any hypervariable DNA sequence loci for use in species-level phylogenetics. CATS-based approaches, as currently implemented, generally

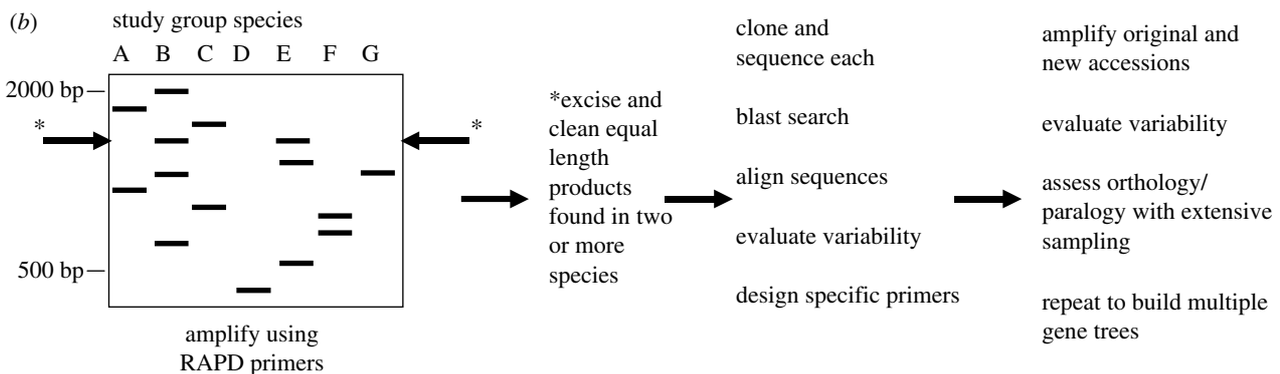
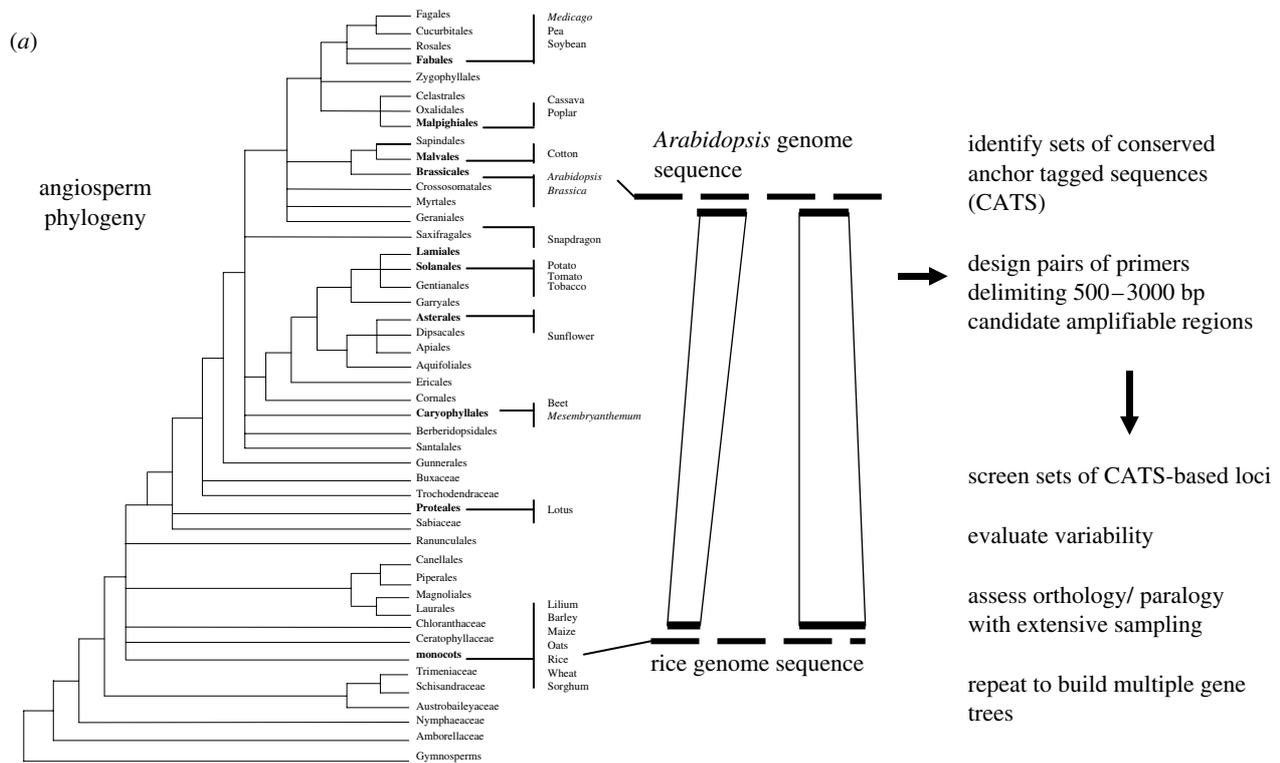


Figure 4. Overview of CATS-based and SCAR-based approaches used to develop nuclear DNA sequence loci. (a) The CATS-based approach, where genomic or EST sequences available for (usually) divergent taxa (phylogeny showing distribution of model taxa for which genomic or EST sequence data are available, modified from Soltis & Soltis 2004), are compared to identify sets of conserved anchor tagged sequences and design primer pairs delimiting 400–3000 bp potentially amplifiable regions that are subsequently screened for variability (Syring *et al.* 2004; Xu *et al.* 2004). (b) The SCAR-based approach where screening for variability in RAPD-generated DNA fragments among closely related species precedes design of primers (modified from Bailey *et al.* 2004).

compare EST/genomic sequences between widely divergent taxa (e.g. rice and *Arabidopsis*; figure 4a). While this may be advantageous for locating widely conserved markers (Fulton *et al.* 2002) thereby facilitating design of potentially ‘universal’ primers, it also means that primers typically span relatively more conserved coding regions. The simple fact that alignments can be constructed between such divergent taxa suggests that such regions are less likely to provide highly variable loci between closely related taxa. However, this is presumably a temporary limitation of our current data bases that include few taxa with many loci while many taxa have very few (or no) loci (e.g. Driskell *et al.* 2004). As the taxon × loci matrix becomes even a bit more densely populated, future CATS-based comparisons among more closely related

taxa will become increasingly powerful for locating DNA sequence loci for specific groups of interest. For example, mapped conifer anchor loci developed from two species of pine (Brown *et al.* 2001) are being used to screen and select potential DNA sequence loci to resolve incongruent and incompletely resolved cpDNA and nrDNA phylogenetic trees for *Pinus* (Syring *et al.* in press). Other examples of comparisons across narrower taxonomic spans illustrate this potential (e.g. in Brassicaceae, Kuittinen *et al.* 2002; Lukens *et al.* 2003 and in monocots, Chandappa *et al.* 2005).

(c) Sequence characterized amplified region-based approaches

Another approach is to use RAPD or AFLP primers to generate and identify sequence characterized amplified

regions (SCARs; Melotto *et al.* 1996) that can be screened as potentially useful sequence loci prior to development of specific primers (Shaw *et al.* 2003; Bailey *et al.* 2004). This strategy is a modified version of the AFLP-based method of McLenachan *et al.* (2000) for characterizing population level gel-based markers. Under this method, PCR products of equal length amplified across a subset of species using commercially available random primers are excised, cloned, sequenced and aligned to evaluate variability (figure 4b; Bailey *et al.* 2004). By using a range of primer combinations many candidate loci can be generated. Specific SCAR primers are then designed for potentially useful regions that show high levels of variability compared to previously used conventional loci such as ITS. As for the CATS-based approaches, the SCAR strategy can be used to screen and generate numerous candidate loci (see below). The SCAR-based approach has been used successfully to increase resolution and support for groups of closely related species of legumes (Bailey *et al.* 2004; see figure 1) and mosses (Shaw *et al.* 2003, 2005a).

There are several potential benefits of using random genomic regions rather than previously characterized genic regions. First, no prior sequence information is required to develop many loci for a group of interest. Second, random amplification of regions across the entire genome eliminates the restriction of working within known genic exon/intron regions (and potentially adjacent 3' UTRs and 5' promoter regions), which are likely to represent the less variable half of the nuclear genome. Given that 53–57% of the *Arabidopsis* genome has been classified as non-coding DNA (Initiative 2000), it presents significant untapped potential as a source of DNA characters (figure 3; Shaw *et al.* 2003, 2005a; Bailey *et al.* 2004). Third, unlike the CATS-based approaches, the initial screening process facilitates selection of loci with levels of variation needed to address species-level problems prior to development of specific primers. Fourth, by homing in on sets of loci with similar high levels of sequence divergence this reduces the chances of locus imbalance where one highly variable locus overrides signal from less variable loci. Finally, by initially selecting for regions with limited length variation this reduces the chances of encountering highly length variable regions that may be difficult to align (e.g. Syring *et al.* in press). This last feature may of course eliminate many potentially useful but length variable DNA sequence loci from consideration (Britten *et al.* 2003; Fondon & Gamer 2004).

A number of potential drawbacks associated with SCAR-based loci are immediately apparent. First, there is no indication that selected loci will be biparentally inherited which is critical for studies of reticulation. BLAST searches can be used to check for significant similarity to complete angiosperm cpDNA or mtDNA sequences available in GenBank, eliminating known uniparentally inherited loci from those sources. Screening via artificial or other known hybrids provides a more certain route to ascertain this. A second potential limitation is the possibility of amplifying regions of junk repeat DNA, with consequent paralogy problems. While this might suggest

greater probability of encountering paralogy problems with SCAR-based loci, in practice all loci are subject to most, if not all of the same considerations. Dealing with paralogy needs to be an integral and central step in screening all DNA sequence loci for phylogenetic analysis (see below). Finally, there are clearly tradeoffs to be weighed up when looking for SCAR-based loci in terms of what span of taxa are to be included in the initial selection of amplified fragments. Inclusion of bands that amplify in just two or three closely related species runs the risk that more divergent taxa will not be amplified by primers developed from the initial sequence data, thereby potentially forfeiting inclusion of outgroup taxa (Bailey *et al.* 2004). Conversely, selecting fragments that amplify across a wider range of taxa than the specific clade of interest may mitigate against finding the hypervariable regions that are needed to resolve relationships within that clade. On the plus side, this is under the control of the investigator; different options can be tested.

5. PILOT STUDIES AND SCREENING

During the short history of plant molecular systematics, the choice of biparentally inherited nuclear loci has been severely limited by the availability of useful primers. The accessibility of new approaches (discussed above) is rapidly expanding the plant systematist's 'toolbox' to a situation with essentially unlimited options (Shaw *et al.* 2003; Bailey *et al.* 2004; Padolina *et al.* 2004; Xu *et al.* 2004; Syring *et al.* in press). For those embarking on the development of new datasets or studies of new groups, these breakthroughs provide tremendous opportunities. To derive maximum benefits from the available loci, investigators need to employ pilot studies incorporating extensive screening to identify the most promising loci prior to investing heavily in any one locus (e.g. Bailey *et al.* 2004; Syring *et al.* in press). The exact order of steps in a screening process will depend on whether or not one is employing an approach that begins with the purchase of specific primers (e.g. CATS as a modified LCNG approach) or the use of random primers (e.g. SCAR-based approaches), but the critical factors that need to be considered remain essentially the same. Any screening approach should consider: (i) how many loci and what DNA samples/taxa to screen; (ii) ease and reliability of PCR amplification and sequencing; (iii) the potential orthology of sequences generated; (iv) alignment difficulties; and (v) last, but certainly not least, relative resolving power of each locus (e.g. Strand *et al.* 1997; Cronn *et al.* 2002; Bailey *et al.* 2004; Syring *et al.* in press).

The number of screened loci and DNA samples used in the selection process are largely dictated by the study group and available resources. Without doubt, selection strategies that screen larger numbers of samples and loci relative to other approaches will provide better starting points. However, these benefits need to be balanced against costs (time and money). The decreasing cost of oligonucleotide synthesis, PCR, and sequencing means that a pre-screening strategy that generates comparative data from 25 to 100 starting loci for 10 or more individuals is generally feasible and

realistic. Such numbers may initially sound excessive, but considering that the goal is to discard problematic loci identified at each level of screening, the actual cost will be much lower than if all loci were carried through every step. Sampling should, whenever possible, focus on non-hybrid diploid individuals spanning the taxonomic breadth of the ultimate study group. Furthermore, the inclusion of some intraspecific samples and samples for which data already exists (e.g. ITS or cpDNA data) will sharpen subsequent screening (e.g. for orthology and variability).

Loci that are easy to amplify and sequence are far preferable to those that are not. Problems with either amplification or sequencing in many accessions suggest that significant unnecessary effort and expense may be involved in primer development, cloning and extra sequencing. Furthermore, templates that are difficult to sequence may be indicative of underlying paralogy problems (see below). Regions which are reasonably straightforward to sequence using PCR primers (plus any additional sequencing primers needed for longer fragments) will facilitate efficient data generation. Of course, there will still be some individual samples that require cloning (e.g. due to hybridization, heterozygosity, or poor primer match), but selection should aim to minimize amplification and sequencing problems.

The assessment of sequence orthology and paralogy can and should be made at several stages. First, simple observation of the number of bands amplified using the 'locus specific' primers can reveal obvious problems. Primer pairs that amplify multiple bands, particularly of similar sizes, should be given lower priority or discarded. Second, difficult to sequence PCR products can be indicative of problems caused by a mixture of paralogous sequencing templates (e.g. Rauscher *et al.* 2002). Furthermore, Scherson *et al.* (in press) have recently noted that repeat patterns of subset polymorphisms in otherwise clean sequencing reads are more likely to be caused by underlying paralogy than by heterozygosity. Consideration of the patterns discussed by Scherson *et al.* (in press) is worthy of inclusion in the screening process. With the large numbers of potential regions now available, there is little reason to intentionally embark on studies that will require the dissection of paralogous gene copies for species-tree reconstruction. Lastly, assessments of orthology/paralogy can be made using comparisons of gene trees developed from each locus in relation to previously generated data (e.g. ITS and cpDNA). In these cases, the inclusion of intraspecific samples can also help identify potential paralogy problems.

Data generated during initial screening to demonstrate reliable single band amplifications and lack of serious sequencing difficulties or obvious paralogy problems can also be used to compare sequence variability, to identify those loci that are easy to use and informative. Assessments of inter-locus variability can be made among the pool of potential new loci and between these loci and previously developed data, such as ITS and cpDNA markers. Comparisons of the former are critical for the selection of the most variable loci, while evaluation of the latter should provide some idea how much additional resolution might (or might not) be gained from a fully sampled matrix using the

'best' of the newly screened loci. Measures of sequence variability may be based on percentage of PI characters, pairwise divergence, or more comprehensive analysis of gene-tree comparisons. Gene-tree approaches best characterize the overall distribution of variability and therefore the relative resolving power of each locus based on the same samples (Wenzel & Siddall 1999). Additional measures of utility for each gene may also be drawn from the individual gene trees (e.g. consistency index, number of resolved nodes, average branch support, etc.).

In plant species-level phylogenetics, rarely has the issue of too much variation been a problem. In general, researchers want to select the most variable loci that fit all the other criteria discussed above. However, one additional measure of variability and potential utility that may be considered in screening is the presence of indels that can provide potentially useful characters, but also potential alignment difficulties. Clean indels contribute useful phylogenetic markers while complicated indel patterns tied to high substitution rates can limit the utility of the data generated. Ideal loci will be relatively straightforward to align while maintaining relatively high levels of information (Bailey *et al.* 2004).

6. SUMMARY COMMENTS

One of the frequently cited advantages of DNA sequence data for reconstructing phylogenies is the virtually unlimited number of characters that can be generated. However, for closely related plant groups it may be extremely difficult to locate sufficiently variable DNA sequence loci to generate enough characters to provide well-resolved and robustly supported hypotheses of species relationships without generating very large volumes of sequence data. In contrast to those who study animals, for whom rapidly evolving mtDNA loci provide a rich source of species-level data, the plant molecular systematist's toolbox lacks readily accessible hypervariable sequence regions. Well-supported resolution at the gene-tree level remains elusive, suggesting limited confidence in how systematists currently infer relationships among closely related species.

Several routes to locating hypervariable nuclear DNA sequence loci for species-level phylogenetic analysis have emerged. The relative efficiency of these alternatives should increase rapidly because locating informative loci depends on how closely related the study group is to species for which significant (genome or EST) DNA sequence is available. SCAR-based approaches seem to present the most efficient strategy for many plant groups at present. However, as the gene \times taxon GenBank matrix become more densely populated, CATS-type approaches based on significantly narrower taxonomic spans will become increasingly frequent and productive.

A number of recent studies have demonstrated the high utility of anonymous, presumed non-coding, non-genic nuclear regions for resolving relationships (figures 1 and 3; Shaw *et al.* 2003, 2005a; Bailey *et al.* 2004). At the same time, it is increasingly clear that although low-copy nuclear introns range greatly in variability (figure 3; Small *et al.* 2004), the majority are

likely to be less variable (at least on a percentage basis) than ITS. Tapping into the large non-genic, or less-conserved genic fractions of plant genomes may thus be particularly productive. At present, the majority of researchers appear to be following CATS-based approaches to develop species-level nuclear DNA sequence loci (e.g. Syring *et al.* in press; Xu *et al.* 2004; Scherson *et al.* in press). In part this may be due to the large number of primer pairs that CATS approaches can provide, or to the attraction of locating universal or at least widely applicable markers. However, we believe that there is a need for caution to avoid the pitfalls associated with the LCNG approach that has been shown to be sub-optimal.

Whether the focus is on LCNG sequences or loci derived from SCAR-based approaches, the solutions will, for the most part, be lineage-specific. Once we stray beyond well-known nrDNA and cpDNA loci, opportunities to develop universal markers or primers appear to be extremely limited, if not non-existent (Sang 2002), suggesting a need to get away from universal gene thinking for species-level phylogenetics. A requirement for marker development, or at least substantive optimization, for specific plant lineages, and potentially for each individual study, shifts the emphasis towards approaches that involve screening prior to primer development. The need to screen candidate loci is further reinforced by the growing ease of identifying large numbers of candidate primer pairs. Thus, whatever sources of sequence data—non-coding cpDNA, LCNG, CATS-based or SCAR-based—are used, strategies for screening candidate sequence loci are key. Given the proliferation of potential sequence loci, screening can afford, and probably needs, to be more ruthless than anything envisaged before.

Finally, none of these developments denies the ongoing utility of, and need for, non-coding cpDNA and ITS data. Not only will these continue to provide useful data in their own right, but cpDNA and ITS data also form essential foundations for subsequent development of other DNA sequence loci. This will take the form of a framework to compare levels of variation, provide initial hypotheses of relationships to direct taxon sampling and assess orthology/paralogy concerns during screening and data generation for candidate nuclear DNA sequence loci.

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GLOSSARY

- AFLP: amplified fragment length polymorphism
 CATS: comparative anchor tagged sequences
 CFI: consensus fork index
 COS: conserved orthologue set
 EMPT: equally most parsimonious tree
 EST: expressed sequence tag
 ITS: internal transcribed spacer
 LCNG: low-copy nuclear gene
 PI: parsimony informative
 RAPD: randomly amplified polymorphic DNA
 SCAR: sequence characterized amplified region
 UTR: untranslated region