

RESEARCH ARTICLES

A Population Genomics Study of the *Arabidopsis* Core Cell Cycle Genes Shows the Signature of Natural Selection ^W

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Large-scale comparison of sequence polymorphism and divergence at numerous genomic loci within and between closely related species can reveal signatures of natural selection. Here, we present a population genomics study based on direct sequencing of 61 mitotic cell cycle genes from 30 *Arabidopsis thaliana* accessions and comparison of the resulting data to the close relative *Arabidopsis lyrata*. We found that the *Arabidopsis* core cell cycle (CCC) machinery is not highly constrained but is subject to different modes of selection. We found patterns of purifying selection for the cyclin-dependent kinase (CDK), CDK subunit, retinoblastoma, and *WEE1* gene families. Other CCC gene families often showed a mix of one or two constrained genes and relaxed purifying selection on the other genes. We found several large effect mutations in *CDKB1;2* that segregate in the species. We found a strong signature of adaptive protein evolution in the Kip-related protein KRP6 and departures from equilibrium at *CDKD;1* and *CYCA3;3* consistent with the operation of selection in these gene regions. Our data suggest that within *Arabidopsis*, the genetic robustness of cell cycle-related processes is more due to functional redundancy than high selective constraint.

INTRODUCTION

The molecular components underlying the mitotic cell cycle show a high degree of functional and structural evolutionary conservation across eukaryotes. However, the transcriptional and posttranslational regulation of the cell cycle can be dramatically different among species (Jensen et al., 2006). This regulatory variation reflects a degree of evolutionary flexibility in the cell cycle that can help explain how plants have evolved several specific cell cycle features, such as lifelong cell divisions leading to new organs or the intrinsic cell totipotency found in most plant species (Gutierrez, 2005; Inzé and De Veylder, 2006). In addition, natural phenotypic variation has been recorded in *Arabidopsis thaliana* for several plant traits related to the cell cycle, such as root growth rate, developmental speed, endoreduplication rate, and shoot size (Beemster et al., 2002; Koornneef et al., 2004; Lisec et al., 2008). However, given the two opposing evolutionary faces of the cell cycle observed between species (conserved in

its components yet highly diverged in its regulation), it is hard to predict a priori if genetic variation present in the cell cycle genes could contribute to natural phenotypic variation within species.

A promising way to evaluate the molecular evolution of cell cycle genes in plants is to incorporate population genetics methods to assess the contribution of positive and negative selection in shaping genetic variation and to determine the overall levels of constraint (Maynard Smith and Haigh, 1974; Nei, 1987; Charlesworth et al., 1993). An added benefit of a population genetics approach is that where evidence of selection is found, it implicitly means that mutations in that locus can result in a change in phenotype that affects the fitness of the individual. Hence, a population genetics approach can equally assist in selecting candidate genes from a molecular system for follow-up studies (Shimizu and Purugganan, 2005; Weigel and Nordborg, 2005; Wright and Gaut, 2005).

A number of population genetic test statistics have been developed to identify single nucleotide polymorphism (SNP) patterns that deviate from expected neutral patterns, both for within-species studies (denoted as polymorphisms) and/or between species (denoted as divergence). A number of neutrality tests contrast polymorphism and divergence patterns in nonsynonymous (NS) and synonymous (S) codon sites (e.g., McDonald and Kreitman, 1991). Other tests make use of the increase (under positive selection) or decrease (under negative selection) in frequency of SNPs or patterns of polymorphism in all sites of a given locus to detect deviations from patterns expected under neutrality (Bamshad and Wooding, 2003; Nielsen, 2005; Biswas and Akey, 2006; Sabeti et al., 2006). Nevertheless, a

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well-recognized problem of these population genetic tests is that non-neutral signatures can also be generated by demographic forces, such as rapid population expansion or population bottlenecks (Tajima, 1989; McDonald and Kreitman, 1991; Eyre-Walker, 2002; Wright and Gaut, 2005). As a consequence, the standard neutral model may not apply for species with complex demographic histories, which can lead to erroneous conclusions about the significance of neutrality test statistics results. However, demographic forces are expected to affect the pattern of polymorphism genome-wide, as opposed to selection, which is expected to act locally on a targeted mutation (Cavalli-Sforza, 1966). One approach to address this issue is to generate an empirical null distribution of test statistic values from a large set of random loci (under the assumption that the vast majority of mutations are essentially neutral; Kimura, 1983). Then, if a test statistic value calculated for a given locus falls in the tails of this null, there is an increased probability that the signature of the locus is a result of selection rather than demography. An alternative approach to differentiate patterns of selection from demography is to model the demographic history of the species and as such estimate the null distribution of the test statistics (Tenaillon et al., 2004; Tenaillon and Tiffin, 2008). However, when the demographic history of a species is complex, a population model that explains all aspects of the data can be elusive. For *A. thaliana*, the demographic models developed so far can only explain parts of the observed data (Nordborg et al., 2005; Schmid et al., 2005; François et al., 2008). Therefore, the empirical approach has been advocated for population genetic analyses in *A. thaliana* (Tenaillon and Tiffin, 2008) and has been successfully applied to reveal potential targets of selection (Kelley et al., 2006; Toomajian et al., 2006; Borevitz et al., 2007).

In this article, we present a study of the mitotic cell cycle in *Arabidopsis* using a population genomics approach. Vandepoele et al. (2002) defined a core set of 61 *A. thaliana* cell cycle genes containing 12 kinases, 30 cyclins, two CDK subunit (CKS) genes, seven Kip-related proteins (KRPs), eight E2F/DP genes, and the Retinoblastoma (*Rb*) and *WEE1* genes. Progression through the four phases of the mitotic cell cycle (first gap, DNA synthesis and replication; second gap, mitosis or abbreviated G1-S-G2-M) is determined by the activity of cyclin-dependent kinases (CDKs) in complex with cyclins (CYCs). The activity of CDK/CYC complexes is induced by CDK-activating kinases and is inhibited in *A. thaliana* by regulators such as KRPs, CKS docking proteins, and *WEE1*. Other cell cycle components are E2F, DP, and *Rb* that form transcription complexes that activate S-phase genes. The complex interplay of these 61 core cell cycle (CCC) and other genes not only leads a cell through mitosis but also initiates the endocycle (when a cell replicates its DNA content without a subsequent mitotic phase) and induces a cell to differentiate (for reviews, see Inzé and De Veylder, 2006; De Veylder et al., 2007).

Here, we describe patterns of sequence variation in the 61 CCC genes in a collection of 30 *A. thaliana* accessions, selected to capture a large fraction of the allelic diversity present in *A. thaliana*. We estimated a series of commonly used population genetics and found evidence of non-neutral processes operating in CCC gene regions using the empirical background distribution approach. For brevity, we refer in this manuscript to the CCC genes as the CCC data set and the empirical background data

set published by Nordborg et al. (2005) as the 2010 data set (Clark et al. 2007). Additional tests of polymorphism and divergence were conducted for the CCC genes using a single *Arabidopsis lyrata* outgroup sequence. The results suggest that only a subset of the CCC genes are subject to strong evolutionary constraints, while others appear to be less constrained or, in some cases, even subject to positive selection.

RESULTS

For all 61 CCC genes we resequenced three or four ~600-bp long fragments in 30 *A. thaliana* accessions to identify SNPs. One fragment was located within 1000-bp upstream of the translation start, one or two fragments were intragenic and covered both intronic and exonic sequence, and one fragment was located within 1000 bp downstream of the translation end. Although we have only partially resequenced the CCC genes, our approach allowed us to assess patterns of variation in the CCC genes. In total, the CCC data set consists of 238 gene fragments from 61 genes in a sample of 30 *A. thaliana* accessions. A complete list of sequences in the CCC data set can be found in Supplemental Data Set 1 online. Of a total of 238 gene fragments resequenced, 99.4% were successful (>300 bp). The average fragment length is 593 bp, and the minimal sample size is 26 accessions. We identified 3329 SNPs in the CCC data set, or one SNP per 42 bp.

Similar to previous findings (Nordborg et al., 2005), we found a strong negative correlation ($R^2 = 0.118$, $P < 0.001$; Pearson correlation) between polymorphism per site (θ_w) in the CCC genes and local gene density (see Supplemental Figure 1 online). No significant differences in exonic ($P = 0.075$), intronic ($P = 0.372$), and downstream ($P = 0.501$) θ_w values were observed between the CCC and the 2010 data sets (Figure 1; see Supplemental Table 2 online). Comparison of θ_w in upstream, exonic, intronic, and downstream site classes showed intronic and downstream θ_w to be similar both within the CCC genes ($P = 0.596$) as within the genes of the 2010 data set ($P = 0.721$). As expected, a significantly lower mean θ_w ($P < 0.001$) in the exonic than in the intronic CCC regions was found, indicating a greater constraint on coding sites. Unexpectedly, we observed a significantly ($P = 0.018$) higher θ_w in the upstream region of the CCC genes compared with the upstream regions in the 2010 data set. The CCC upstream θ_w value was also significantly higher than the CCC intronic and downstream θ_w ($P = 0.002$ and 0.017 , respectively). These differences between upstream and intragenic/downstream θ_w values were absent for the 2010 data set. Mutation frequencies positively correlate with GC content (Hurst and Williams, 2000), so a higher GC% could explain θ_w differences. However, we found GC% in all four regions to be higher in the 2010 data set than in the CCC data set. Within the CCC data set, GC% in the upstream region (31.6%) was similar to GC% in intronic and downstream regions (32.4 and 31.5%, respectively). This is an unusually high level of nucleotide polymorphism in upstream gene regions. The difference may be explained by an unequal presence of functional promoter motifs in the upstream fragments of both data sets. It is also possible that what we observe is an excess of slightly deleterious mutations in the CCC upstream regions as a consequence of reduced selective constraint (Bustamante et al., 2002; Foxe et al., 2008).

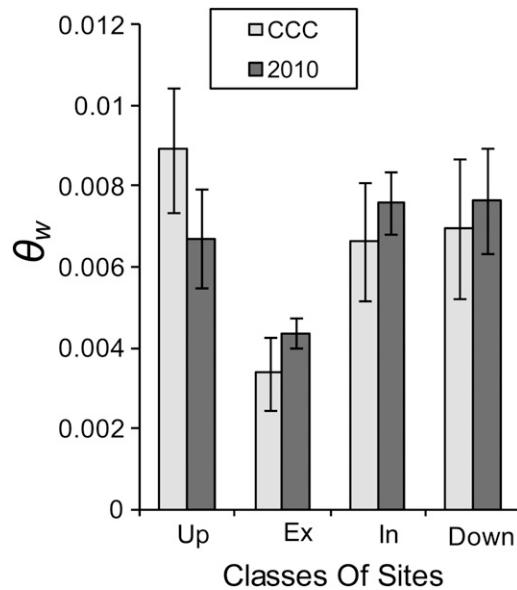


Figure 1. Histogram of the Average θ_w Values in the Upstream, Exonic, Intronic, and Downstream Sites Analyzed in the CCC Data Set and the 2010 Data Set.

Bars represent the 95% confidence interval. See Supplemental Table 2 online.

Patterns of SNP Diversity in CCC Coding Sequence

When mutations occur in the coding sequence of a gene, this can lead to amino acid or gene integrity changes, such as altered splice sites or premature stop codons. Such mutations can result in important alterations of protein function and may therefore be subject to natural selection. For *CYCD7;1*, *CYCA3;3*, and *CDKB1;2*, we found polymorphisms in the coding sequence that introduced premature stop codons in some accessions relative to the Col-0 reference (see Supplemental Table 3 online). The stop codons introduced in *CYCD7;1* and *CYCA3;3* were located at the C-terminal end of the amino acid sequence. In *CDKB1;2*,

we identified three haplotypes in which a single base deletion resulted in early stop codons that truncated >50% of the protein sequence (see Supplemental Table 3 online). This is surprising because plant CDK sequences consist largely of functional domains (Joubès et al., 2000). In addition, we found that the *A. lyrata* ortholog of *CDKB1;2* lacked most of the first two exons of the *A. thaliana* copy, equal to 70% of this protein (Figure 2), which we reconfirmed by PCR amplification and sequencing. However, we did not observe large-effect SNPs in the paralogous *CDKB1;1* sequence. These observations suggest that *CDKB1;2* is not essential for the fitness of the plant and may be on its way out of the *Arabidopsis* gene pool. Because of its disrupted integrity, *CDKB1;2* was excluded from further coding sequence diversity analysis.

Patterns of molecular evolution in CCC genes can be assessed with comparisons of the ratio of NS changes per NS site to the ratio of S changes per S site. We abbreviate these ratios as pN/pS for polymorphism and Ka/Ks for divergence. Because amino acid polymorphisms in the coding regions were not abundant enough to evaluate the pN/pS ratio at individual genes, the mean pN/pS for groups of CCC genes was used to evaluate the level of purifying selection within *A. thaliana* (Table 1). Overall, we found an average pN/pS of 0.304 in the 61 CCC genes, which was significantly higher ($P < 0.05$) than the genome-wide average in the 2010 data set ($pN/pS = 0.252$). The CDK family had a pN/pS value of 0.119, which is less than half of any other CCC gene family ($P < 0.05$). The high pN/pS value for cyclins is partly due to two outliers; *CYCA3;3* has 24 NS to one S and *CYCB1;3* has eight NS and no synonymous polymorphisms. Excluding these outliers lowered pN/pS for cyclins to 0.308, which is still high compared with the genome-wide average of 0.252 ($P < 0.05$).

The Ka/Ks ratio of each CCC gene (see Supplemental Table 4 online) was estimated on the full reference coding sequence and clustered with the partitioning around medoids (PAM) algorithm (Kaufman and Rousseeuw, 1990). Clustering of Ka/Ks was done to illustrate different degrees of constraint across the cell cycle machinery. Wilks' Lambda test statistic suggested six clusters as optimal partitioning (Figure 3). We noticed that the cluster with the lowest mean Ka/Ks (cluster 1) contained all but one CDK (*CDKF;1*; cluster 2). Interestingly, the tandem duplicated genes

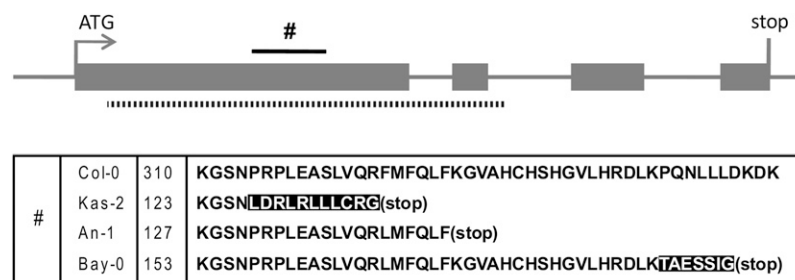


Figure 2. Schematic Representation of *CDKB1;2* in *Arabidopsis* and the Large-Effect Changes Observed.

Blocks in the figure represent exons annotated in *A. thaliana*. The dashed line delimits the gene region that was deleted in the *A. lyrata* orthologous *CDKB1;2* gene. The solid line (#) delimits the amino acid region where early stop codons were observed in *A. thaliana*. Changes in the amino acid sequence within *A. thaliana* are summarized. The numbers are the position of the stop codon in the amino acid sequence. Each amino acid sequence represents a haplotype, symbolized by one accession. The Columbia-0 sequence is the reference. Changes in amino acid sequence due to a frame shift are marked in black.

Table 1. Estimates of the Mean pN/pS and Ka/Ks Ratios

CCC Group	Mean pN/pS	Mean Ka/Ks
CCC Gene Family		
CDKs	0.119 ^a	0.073 ^a
Cyclins	0.357 (0.308 ^b)	0.241
E2f/DP	0.324	0.239
Interactors	0.294	0.300
CCC PAM cluster		
Cluster 1	0.132	0.065
Cluster 2	0.239	0.156
Cluster 3	0.327	0.223
Cluster 4	0.410	0.290
Cluster 5	1.230	0.413
Cluster 6	0.270	0.597

^aWithout CDKB1;2.^bWithout CYCA3;3 and CYCB1;3.

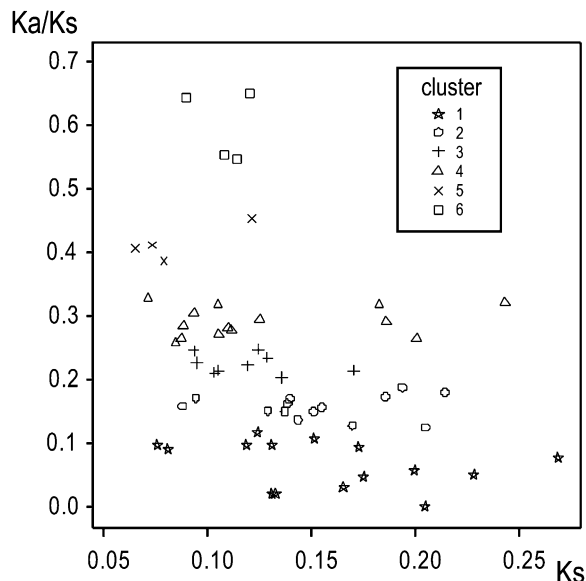
CKS1 and *CKS2* were also contained in cluster 1. The reported differential expression pattern of the CKS genes (Menges et al., 2005) and the high selective constraint suggests that neo- or subfunctionalization happened after the duplication event, which was prior to the divergence of the *Arabidopsis* lineage. Also, cluster 2, which contained the single-copy genes *Rb* and *WEE1*, showed high levels of selective constraint. Because the rate of evolution has been found to negatively correlate with gene expression (Wright et al., 2004; Foxe et al., 2008), we examined the effects of CCC expression levels (Menges et al., 2005) on the evolutionary rate by analysis of variance. We found that cluster 1 shows significant ($P < 0.05$) higher gene expression levels across all reported plant tissues compared with the other clusters. For all gene families other than the CDKs and CKSs, we saw different rates of evolution across the genes within the family. Some clear examples were the DEL and KRP families. The Ka/Ks ratios of the three DEL family genes were 0.123, 0.268, and 0.643. *KRP2* and *KRP3* were found in cluster 2, while *KRP1*, *KRP5*, *KRP6*, and *KRP7* were all found in cluster 5 or 6. The variation in evolutionary rate might indicate which members of the family play a more evolutionary conserved role in the overall cell cycle machinery and which genes are more likely to be subject to adaptive change.

We compared Ka/Ks and pN/pS values calculated for the six PAM clusters and for the four major functional groups of the CCC genes (Table 1). PAM cluster 1 and the CDK family had both the lowest pN/pS and Ka/Ks value. This suggests that purifying selection against NS polymorphisms within species mirrored the observed slow rate of evolution in between-species comparisons for these functional groups. Less concordance between pN/pS and Ka/Ks values was found in the other groups.

The MK test is based on the neutral expectation that the ratio of NS to S mutations will be equivalent for intraspecific polymorphism and between-species divergence (McDonald and Kreitman, 1991). The direction and average strength of selection on AA changes was estimated by the MKPRF selection parameter γ . Selection against NS replacements results in $\gamma < 0$, selection in favor of NS replacement in $\gamma > 0$. The MK test after Bonferroni correction was significant in *CYCA3;3* and *CYCB1;3*

($P < 0.05$) in the direction of an excess of NS polymorphisms. This excess of replacement polymorphism is consistent with selective processes, including a recent relaxation of functional constraints, segregation of slightly deleterious mutations, local adaptation, or some form of balancing selection. The results for MKPRF similarly suggest that purifying selection may be a primary form of selection on amino acid changes in this set of genes. Estimates of the selection parameter (γ) were significantly negative for *CYCB2;1*, *CYCA3;1*, *CYCD7;1*, *CYCA3;3*, and *CDKD;1* but not for *CYCB1;3* (see Supplemental Table 4 online). Many similar observations have been made for a variety of genes in *Arabidopsis* and have consistently been interpreted as evidence for an excess of slightly deleterious variation segregating in the *Arabidopsis* genome (Bustamante et al., 2002; Foxe et al., 2008). However, for *CYCA3;3*, *CDKD;1*, and *CYCB2;1*, additional signatures for departures from neutrality were observed in the genomic region of these genes (see below).

For one gene, *KRP6*, the MK test was significant ($P < 0.05$) due to an excess of replacement divergence relative to polymorphism, although this was not significant after Bonferroni correction. *KRP6* also had the highest estimated selection parameter (γ) of all CCC genes, although again not significant. Because MK and MKPRF tested for selection from the resequenced *KRP6* fragments only (for *KRP6* this equals 80% of the total reference coding sequence), we estimated Ka/Ks again on the resequenced part of *KRP6*. For this section of the gene, we found Ka/Ks to be 1.10, which also is consistent with adaptive substitution. We estimated the 95% Ka/Ks confidence interval for *KRP6* from 0.3955 to 3.8685 (Comeron 1995), implying that we cannot strictly reject that Ka/Ks equals one. However, the only

**Figure 3.** Scatterplot of the Ka/Ks versus Ks Values per Gene.

Ka/Ks values are PAM clustered into six clusters and made visually distinguishable. Dotted lines were added to clarify the six clusters boundaries. Each marker represents one CCC gene. The gene content of each cluster is summarized in Supplemental Table 4 online.

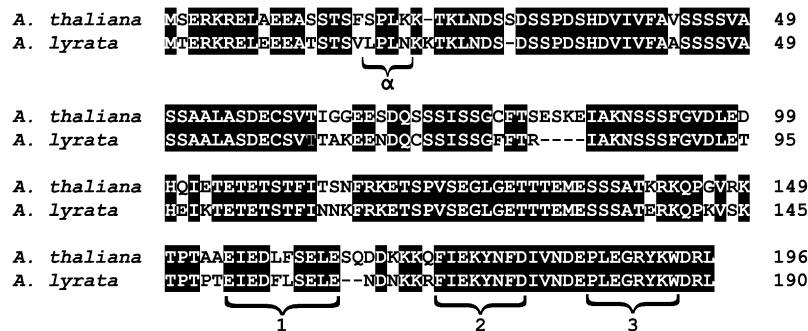


Figure 4. Protein Sequence Alignment of *KRP6*.

The position of the CDK consensus phosphorylation site is indicated by α , and the positions of the conserved amino acid motifs in plant KRPs are indicated by 1, 2, and 3.

reported genes with similarly high Ka/Ks ratios are a very few cases of *R* genes (Xiao et al., 2004; Bakker et al., 2006). Closer inspection of the *KRP6* residues that have fixed since divergence with *A. lyrata* indicated that two replacement fixations were located in a conserved C-terminal CDK and cyclin binding motif region (Figure 4). Also, the N-terminal CDK phosphorylation site, which has been suggested to determine the stability and temporal degradation of the protein (De Veylder et al., 2001), showed two replacement fixations. These results are consistent with a scenario of adaptive evolution of *KRP6* since the divergence of the two *Arabidopsis* species.

Patterns of SNP Diversity across the CCC Sequence Loci

Positive selection on a locus can result in patterns of polymorphism and divergence that depart from neutral patterns driven by mutation and genetic drift. Frequency spectrum-based statistics provide one mean of quantifying these patterns and can detect departures from equilibrium generated by two major types of positive selection. A selective sweep (when a beneficial allele becomes fixed in a population) is hallmarked by an excess of rare SNPs within a region of overall low nucleotide diversity. Balancing selection (when more than one allele of the gene is maintained in a population) is characterized by an excess of intermediate frequency SNPs in a region of high nucleotide diversity (Hudson and Kaplan, 1988; Bamshad and Wooding, 2003; Nielsen, 2005; Biswas and Akey, 2006; Walsh, 2008).

Departures from neutrality at the CCC loci can be tested with the frequency-spectrum based statistics Tajima's *D* (Tajima, 1989) and Fu and Li's D^* and F^* (Fu and Li, 1993) or with comparable tests Fu and Li's D and F (Fu and Li, 1993) and Fay and Wu's *H* (Fay and Wu, 2000) that require an outgroup (see Supplemental Data Set 2 online). In this article, we will only discuss Tajima's *D* statistics and refer the reader to Supplemental Data Set 2 online for the Fu and Li statistics. Significant positive values for Tajima's *D* statistics are consistent with the pattern of a locus under balancing selection. Significantly negative values suggest that the locus has been subjected to a selective sweep. A significantly positive Fay and Wu's *H* indicates a deficit of moderate- and high-frequency derived SNPs

relative to equilibrium expectations, whereas a significant negative Fay and Wu's *H* indicates an excess of high-frequency derived SNPs. We assessed potential significance of Tajima's *D* using the empirical approach and weighed Tajima's *D* values against the genome-wide distribution of Tajima's *D* for the selected 30 accessions, as inferred from the 2010 data. A potential problem for the empirical approach is that the fragments for each CCC gene in our data sets are linked within a 5-kb window. The distribution of Tajima's *D* calculated on the CCC data set was similar to the distribution of the 2010 data set (two-sample smooth test: $P = 0.990$; Figure 5). We observed an unexpected small increase in positive values for Tajima's *D* (Figure 5), but this was not significant (skewness component of two sample smooth test: $P = 0.626$, F-test on distribution variance: $P = 0.76$). We did not have genome-wide outgroup sequence data at hand to calculate the genome-wide distribution

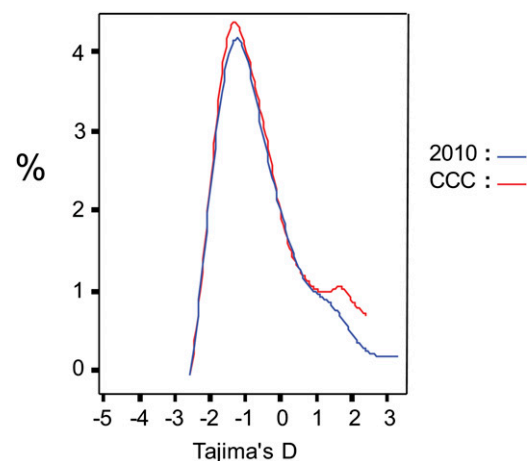


Figure 5. Estimated Genome-Wide Distribution Plots for Tajima's *D* Neutrality Test Statistics.

Red lines represent the test statistic values calculated for 234 CCC gene fragments and blue lines the empirical genome-wide distribution calculated based on 973 fragments from the 2010 data set. The y axis indicates the frequency of the test statistic value in the data set.

of Fay and Wu's H. To avoid confusion, we report significance for Tajima's D values that exceed the negative or positive 2.5% thresholds of the 2010 genome wide distribution but report percentiles of the CCC distribution for the Fay and Wu's H values.

Many CCC genes showed outlier test statistic values, yet we observed high variation in nucleotide diversity levels and frequency patterns between the resequenced fragments even within the same CCC gene. For example, only 32 CCC genes have similar Tajima's D values (coefficient of variation range from -1 to 1) across all the resequenced fragments in the particular gene region. Such high variation has been reported before in *A. thaliana* (Tian et al., 2002; Shepard and Purugganan, 2003; Ding et al., 2007; Moore and Stevens, 2008), necessitating a conservative approach when interpreting population genetics statistics. The unusual patterns of variation in *CDKD1*, *CYCA3;3*, and *CYCB2;1* seemed most suggestive of a departure from neutrality due to selection on these loci or closely linked loci. *CDKD1* showed high negative Tajima's D values in all four fragments and was significant in two. Fay and Wu's H was also negative in all four fragments in the region, with the most extreme value of -19.5 in the first intragenic fragment (98th percentile). *CYCA3;3* showed a very similar pattern with negative test statistics throughout the region. Examination of the derived site frequency spectrum showed a U-shaped pattern with SNPs limited to the lowest and highest frequency classes for both genes (Figure 6). The high frequency-derived polymorphism patterns were driven by the presence of a single haplotype that harbored mostly ancestral variation (represented by Ga-0 for *CDKD1* [Figure 7A] and Ei-2 for *CYCA3;3* [Figure 7B]).

CYCB2;1 showed a very complex pattern of variation, with significant positive Tajima's D values in the upstream region and a highly negative Fay and Wu's H value (99th percentile) for the first intragenic fragment. This signature suggests balancing selection; however, the frequencies of the different haplotype blocks varied between the resequenced regions, most probably as a result of intragenic recombination (Figure 7C). We also observed many derived polymorphisms both at high and at intermediate frequencies (Figure 6). Despite being an interesting pattern (especially given the significantly negative selection parameter γ for *CYCB2;1*; see Supplemental Table 4 online), which has been observed and linked to balancing selection before (Innan et al., 1996; McDowell et al., 1998), it is hard to draw any clear conclusions on the history of selection for this locus.

DISCUSSION

In summary, we have found that the patterns of DNA sequence variation in the 61 CCC genes studied here suggest that within *Arabidopsis* the molecular cell cycle machinery is not a rigorously conserved system but that it is subject to different types of selection. When comparing divergence between *A. thaliana* and *A. lyrata*, all CCC genes showed $Ka/Ks < 1$, yet there is a high degree of variation in the rate of protein evolution across this system of genes. As expected, we found that many CCC genes showed evidence of strong selection against amino acid divergence (e.g., across the CDK family). Highly constrained genes are believed to occupy the vital positions in the molecular

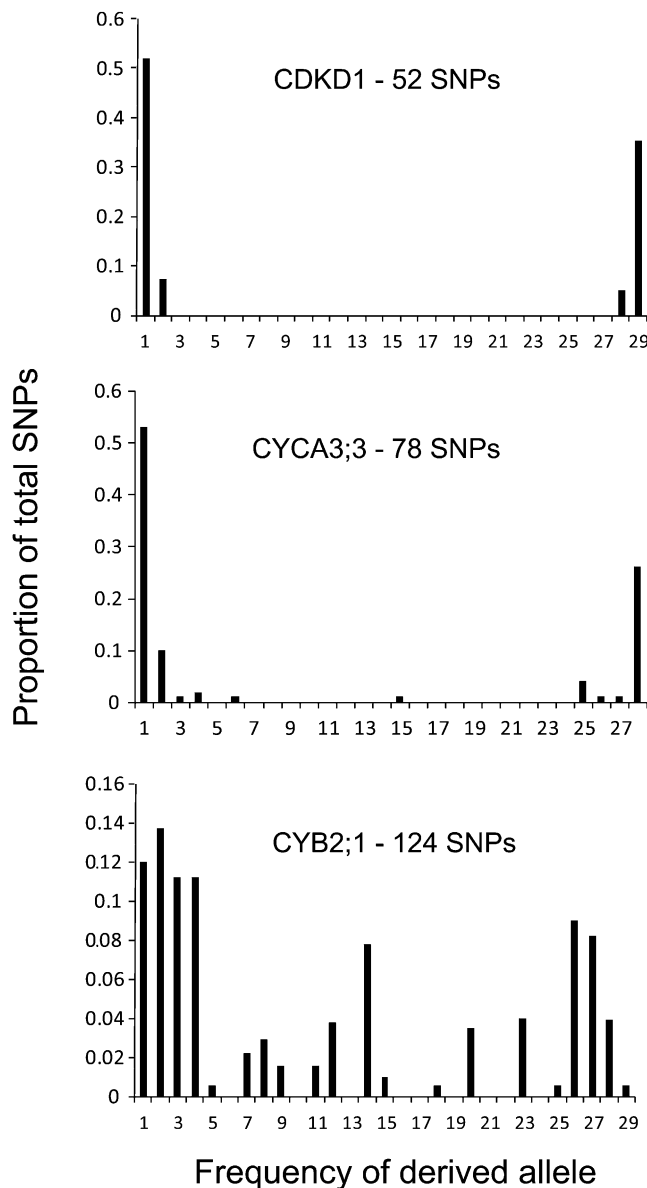


Figure 6. Derived Site Frequency Spectra for *CDKD1*, *CYCA3;3*, and *CYCB2;1*.

The x axis represents the count frequency of the derived SNP allele within 30 accessions. The y axis represents the frequency of SNP sites that have the given derived SNP frequency of the x axis. Bay-0 was removed from the *CYCA3;3* analysis because of missing data.

network, as has been demonstrated for some CCC genes: loss-of-function mutations in *Rb* or *CDKA1* lead to gametophytic lethality (Ebel et al., 2004; Iwakawa et al., 2006; Nowack et al., 2006), and overexpressing *WEE1* arrests the cell cycle progression (De Schutter et al., 2007). However, we also found high rates of amino acid substitution, suggesting fundamental differences in levels of selective constraint across different CCC gene families. Most unexpected was the observation that the mean pN/pS ratio in the CCC genes is higher than in genes

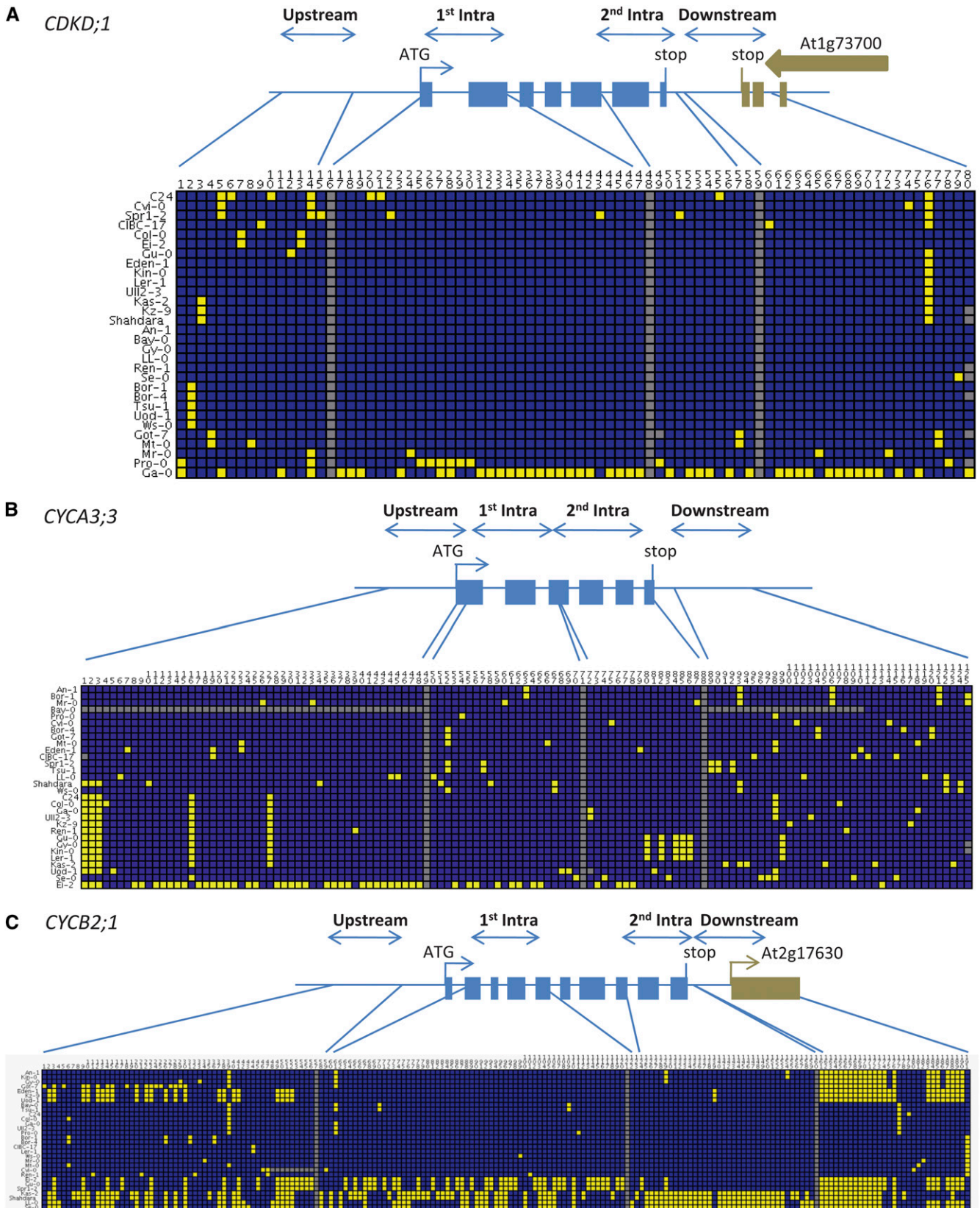


Figure 7. Visual Representation of Haplotypes of the *CDKD;1*, *CYCA3;3*, and *CYCB2;1* Loci.

Each row represents an accession and each column a polymorphic site. For each polymorphic site, minor alleles are in yellow, major alleles in blue, and missing data in gray. Accessions are clustered by haplotype. Gray columns separate the resequenced fragments on the locus. On the schematic gene representation, blocks represent exons.

selected randomly, again suggesting lower selective constraints on many cell cycle genes despite their posited functional importance.

This variation in selective constraint of the CCC genes has several potential origins. One explanation is that gene expression negatively correlates with the rate of protein evolution in *Arabidopsis* (Wright et al., 2004; Foxe et al., 2008), and differences in expression level could therefore account for this pattern. Consistent with this hypothesis, many of the CCC genes where Ka/Ks was low have the highest expression among CCC genes across different plant tissues (Menges et al., 2005). Less-constrained CCC genes have low expression or only high temporal expression levels in specific tissues (Menges et al., 2005). Another possible contributing factor to the observed differences in selective constraints might be related to differences in the ratio of functional domains relative to the size of the protein. Kinases have more functional domains (Joubès et al., 2000) relative to their size than KRPs (De Veylder et al., 2001) or cyclins (Wang et al., 2004). Third, differences in the evolutionary rates may also reflect a difference in the degree of functional redundancy within the different families. Functional redundancy is usually, but not exclusively, a consequence of gene duplication (Pickett and Meeks-Wagner, 1995). If two genes are redundant, this can lead to a decrease in selective constraint on one or both genes (Ohno, 1970). However, selective constraint to maintain this redundancy has also been reported (Moore and Purugganan, 2005). The core cell cycle machinery contains many duplicated genes (Vandepoele et al., 2002), and functional redundancy has been reported for CYCA2-type genes (Imai et al., 2006) and knockout studies of the CDKB1 type (Boudolf et al., 2004). In our data set, we see examples of increased, as well as decreased, rates of evolution for duplicated and functionally redundant genes. The tandem duplicated CKS genes showed Ka/Ks values indicative for strong constraint, while for the *CDKB1;2* gene, integrity changes lead to molecular nonfunctionalization in both *A. thaliana* and *A. lyrata*. This apparent nonfunctionalization of *CDKB1;2* in both *Arabidopsis* species could be a consequence of functional redundancy with *CDKB1;1* (Boudolf et al., 2004). However, *CDKB1;2* seems to be the only CCC gene where nonfunctionalization is as clear despite the high level of duplication and redundancy in the rest of the cell cycle.

The model that emerges from these observations suggests that the robustness of plant development and cell cycle progression is secured by functional redundancy rather than by selective constraint against amino acid substitution. Under this scenario, changes in the timing and mode of expression are more likely to have an impact on plant physiology and therefore drive functional diversification in the cell cycle (Blanc and Wolfe, 2004; Menges et al., 2005; Jensen et al., 2006). Interesting in that regard, yet speculative, is the observed significantly increased θ_w in the upstream fragments of the CCC genes relative to the genome-wide upstream θ_w . This may be the result of differences in the density of functional promoter sites between both data sets. Alternatively, variation in purifying selection between the CCC genes and the 2010 data sets could explain our observation. The decrease in selective constraint on the CCC promoters would allow an increase in CCC regulatory variation across *A. thaliana* populations and drive the evolution of the cell cycle, as demonstrated by Jensen et al. (2006).

In addition to the observation that many CCC genes are not subject to strong selective constraints, we compiled evidence to suggest that some CCC genes harbor molecular patterns consistent with the operation of positive selection. *KRP6* showed a Ka/Ks ratio of 1.1 and had fixed mutations in known functional domains. Furthermore, *KRP6* showed a significant MK test ($P < 0.05$) due to an excess in replacement divergence and had the highest positive MKPRF selection parameter (γ) of all CCC genes. This excludes the possibility of relaxed purifying selection for this gene in *A. thaliana*. The high Ka/Ks value together with the MK test is consistent with historical positive selection operating on this gene. It is therefore interesting that *KRP6* is the only KRP to have transcriptional activation at the S-to-G2 transition in the cell cycle (Menges et al., 2005) and so far the only KRP identified to be degraded at the protein level at the spindle assembly checkpoint during mitosis (G. De Jaeger, personal communication).

Two other CCC genes, *CDKD;1* and *CYCA3;3*, showed negative values for the frequency spectrum-based statistics and a U-shaped derived site frequency spectrum. Demographic scenarios are capable of generating a U-shaped frequency distribution (Caicedo et al., 2007). However, we find demography alone to be an unlikely explanation because for Ga-0 and Ei-2, we observed only one CCC locus with an ancestral haplotype outlier in these accessions. Under a demographic scenario, we would see haplotype outliers in Ga-0 and Ei-0 over multiple loci across the genome, as observed in the geographical isolates Cvi-0 and Mr-0 (Nordborg et al., 2005; François et al., 2008). However, if high variability of unusual coalescents in the genome of this predominantly selfing species exists, almost any pattern, including the pattern we observed, could be generated at any locus in the genome, and a demographic scenario cannot be excluded. Both genes also had a significantly negative selection parameter γ in MKPRF because of an excess of amino acid polymorphism. A negative γ is consistent with purifying selection on slightly deleterious mutations, but for *CDKD;1*, we found five out of seven amino acid mutations in Ga-0 and for *CYCA3;3* 13 out of 24 amino acid mutations in Ei-2. Possible selective explanations for the negative γ and frequency statistic values are balancing selection (with the selective signature being masked by the sample strategy), local adaptation, haplotype introgression, or that the genes are located in a region that underwent a partial selective sweep.

A few interesting biological features for *CDKD;1* and *CYCA3;3* provide a working hypothesis on how natural variation in these genes would affect fitness. Increased expression of the *Oryza sativa CDKD;1* accelerated S-phase progression and cell growth rates (Fabian-Marwedel et al., 2002). From this result it is thought that cyclin activating kinases, like *CDKD;1*, can play an important role in determining the growth rate and differentiation status of cells by controlling CDK activity. However, knockout studies of *CDKD;1* in *A. thaliana* showed no change in phenotype under normal growth conditions (Shimotohno et al., 2006), suggesting functional redundancy with other CDKs. *CYCA3;3* is the only A-type cyclin without a Destruction-box protein domain (needed for rapid degradation of cyclins to terminate the activity of the CDK/CYC complex; Wang et al., 2004). The gene also shows a constant expression profile during the cell cycle, which is

atypical for A3-type cyclins (Menges et al., 2005), but higher expression specifically in anthers (Wang et al., 2004). This could suggest that neofunctionalization of this cyclin after duplication shifted it out of the mitotic cell cycle into the pathway for floral organ differentiation.

The plant cell cycle system coordinates many essential cell fate decisions, such as cell differentiation, cell proliferation, mitotic arrest under stress, and endoreduplication. It was therefore somewhat surprising that the cell cycle machinery, where conservation was expected, displayed a variety in signatures of natural selection. This evolutionary flexibility in the genes driving the cell cycle might contribute to a plant's ability to adapt to environmental changes. The population genomic approach demonstrated in this article contributes to a better understanding of how the cell cycle evolves and has the additional value to assist in nominating candidate genes for extended genetic studies. The CCC genes that display interesting allelic variation, in particular in combination with signals of positive selection, are prime candidate genes for follow-up studies, such as linkage and association studies, ascertaining the effect of cloned CCC alleles across different accessions or cloning *A. lyrata* paralogous genes in an *A. thaliana* background.

METHODS

Selected Accessions

The sample of 30 *Arabidopsis thaliana* accessions was selected from the collection of 96 accessions described by Nordborg et al. (2005) (Nottingham Arabidopsis Stock Centre stock number N22660). A set of 11 accessions was chosen based on their use in generating mapping populations (An-1, C24, Col-0, Cvi-0, Gu-0, Gy-0, Kas-2, Ler-1, Shahdara, Tsu-1, and Ws-0). This initial core of accessions was further enriched for allelic variation through the M strategy of Schoen and Brown (1993) as implemented in the software algorithm *Mstrat v1.0* (Gouesnard et al., 2001). The M strategy selects a set of 30 accessions that captures almost all of the allelic diversity present in the set of 96 accessions (McKhann et al., 2004). We ran *Mstrat* on 25 polymorphic fragments from the 2010 data set, evenly dispersed over the genome and in linkage equilibrium. Using 100 iterations per *Mstrat* run (with Nei's diversity index [Nei, 1987] as second criterion of maximization and the 11 predefined accessions were set as kernel core), we constructed 300 independent core sets of 30 accessions and retained the 30 most selected ones. The final core collection (see Supplemental Table 1 online) successfully picked up accessions from every geographic subpopulation identified by Nordborg et al. (2005).

DNA Samples and Sequencing

The genes studied were the 61 core cell cycle genes of *A. thaliana* as defined by Vandepoele et al. (2002). For *A. thaliana*, we resequenced four ~600-bp fragments per CCC gene positioned as follows: one fragment in the 1000-bp upstream sequence from the translation start, one intragenic fragment covering both exonic and intronic DNA close to the translation start, one intragenic fragment located near the 3' end of the gene, and one fragment in the 1000-bp sequence downstream of the translation end. For the 2010 data set, we used the same 1000-bp window criterion to classify sites as up- or downstream. For the CCC data set, we used the most recent gene models and 1000-bp up- and downstream sequences from the Arabidopsis Genome Initiative reference sequence available on The Arabidopsis Information Resource at the moment of analysis (TAIR6;

<http://www.Arabidopsis.org>). The choice to resequence well-positioned fragments instead of the whole gene region was to keep resequencing costs and efforts at a minimum, while still generating a complete overview of the nucleotide diversity in each cell cycle gene and its direct surroundings. Only one intragenic fragment was designed in *CKS1*, *CKS2*, *KRP5*, and *KRP7* because of their small gene size. All primers were designed with the *PRIMER3* software (Rozen and Skaletsky, 2000). Primers were chosen without regard to predicted functional regions. Genomic DNA was extracted from the shoot tissues with the CTAB method (Doyle and Doyle, 1990). Amplification was done using PCR protocols for direct sequencing with SilverStar DNA polymerase from Eurogentec. DNA fragments were purified with ExoSAP-IT (USB) and sequenced directly in forward orientation using cycle sequencing with the BigDye Terminator v3.1 kit (Applied Biosystems) on an ABI-3000 automated sequencer. The resequencing of the *CDKF1* upstream region recurrently failed for unclear reasons. We found the upstream region of the unrelated gene At4g09390 to be very similar to the *CDKF1* upstream region and suggest the problem to be coamplification with this region. From the downstream sequence of *KRP2*, only 52 bp was resequenced because of poly-T regions in the fragment. A complete list of sequences in the CCC data set and the primers used can be found in Supplemental Data Set 1 online. An 8x coverage genome assembly of *Arabidopsis lyrata* subsp *lyrata* was provided to us by the Department of Energy Joint Genome Initiative *A. lyrata* genome sequencing project. The *A. lyrata* sequence trace archives are accessible at the National Center for Biotechnology Information (<http://www.ncbi.nlm.nih.gov/>). Orthologous *A. lyrata* CCC genes were localized with BLAST software (Altschul et al., 1990) and locally aligned with *A. thaliana* with the Smith-Waterman algorithm (Smith and Waterman, 1981).

SNP Detection

Base calling, sequence assembly, sequence alignment, and polymorphism identification were performed with the *Phred/Phrap/Consed* package (Ewing et al., 1998; Gordon et al., 1998). All sequences and polymorphisms were validated by visual inspection of the chromatograms and edited where appropriate. In case of low-quality sequences (*Phred* quality score <20) or ambiguous SNP identification, the whole fragment was again resequenced in both forward and reverse orientations. Heterozygous SNPs and sequences whose quality remained low despite multiple attempts were removed from further analysis. Processing and archiving in multi-fasta files was done with the MEGA3 version 3.1 software package (Kumar et al., 2004). Haplotype visualization was done with the online VH1 software package (<http://pga.gs.washington.edu/VH1.html>).

Population Genetic Analysis

All analyses within or between the CCC and 2010 data sets were done on the 30 selected *A. thaliana* accessions only. Sites with insertions, deletions, or with more than two SNP alleles were discarded from all population genetic analyses because of the complex mutational patterns. All estimates of local nucleotide diversity θ_w are based on the number SNPs per site (Watterson, 1975) and were calculated with Nei's θ_w estimation (normalized θ_w for analyzed sites per locus; Nei, 1987; equation 10.3, $n = 30$). θ_w calculations in up- and downstream fragments were done on noncoding sites only. Local gene densities surrounding the CCC genes were retrieved from Nordborg et al. (2005), and correlations with θ_w were calculated with Pearson's correlation. θ_w of a fragment or of a class of sites (e.g., synonymous) was only calculated when the number of resequenced sites was at least 100 bp. For calculations of θ_w , θ_N , and θ_S in exons, a minimum of 240-bp exonic sites per fragment was applied to retain sufficient amounts of synonymous sites per fragment. Assessment of the significance of differences in mean θ_w values between classes of

sites within or between the CCC and 2010 data sets was done with one-way analysis of variance on log-normalized θ_w values.

Estimation of exonic S and NS sites, pN/pS and Ka/Ks values, and calculating MK tests were conducted with polydNdS, Gestimator, and MKtest in the libsequence software package (Thornton, 2003). pN/pS comparisons were tested for significance using χ^2 on the pooled NS and S counts per tested proportion. Ka/Ks was estimated using the method of Comeron (1995) and tested for significance using the K-estimator software package (Comeron, 1999). Ka/Ks estimates are based on the full coding sequence in the Columbia-0 and *A. lyrata* comparison. Ka/Ks was clustered using the PAM algorithm (Kaufman and Rousseeuw, 1990) using the R environment for statistical computing (<http://www.r-project.org/>). PAM searches for k representative objects or medoids for the values in the data set. Then, k clusters are constructed by assigning each observation to the nearest medoid. Optimization of k was done with Wilks' Lambda test. MK tests and MKPRF analysis were done on coding sites from the resequenced CCC fragments only. Significance of the MK test was Bonferroni corrected for multiple testing. The population selection parameter γ and its 95% credibility interval were estimated using Markov chain Monte Carlo algorithms and the default settings of the priors on the MKPRF web server (<http://cbsuapps.tx.cornell.edu/mkprf.aspx>), which is partially funded by Microsoft. The MKPRF algorithm scales the intensity of γ for every individual gene to the selection intensity distribution observed for all genes using a Bayesian implementation of Poisson random field (Sawyer and Hartl, 1992; Bustamante et al., 2002), and estimates of γ represent the mean of the posterior distribution. Because MKPRF tests NS/S counts against the expected model derived from all genes tested, it is considered more powerful to detect selection than MK (Bustamante et al., 2002) but has shown to have a linear dependency between the number of detected genes with $\gamma > 0$ and the priors used for the model (Li et al., 2008).

Neutrality test statistics Tajima's D (Tajima, 1989), Fu and Li's D/D^* and F/F^* (Fu and Li, 1993), and Fay and Wu's H (Fay and Wu, 2000) were calculated using the software VariScan v2.0 (Hutter et al., 2006). The neutrality test statistic formulas in Variscan use Nei's θ_w estimator and a modification of the pairwise nucleotide diversity statistic θ_π (Tajima, 1989), π_m , which can handle missing data at sites without the need of discarding them (Hutter et al., 2006). All test statistics calculations were done with the Variscan parameters set to only analyze sites with <10% missing data (referred to as informative sites). In both the CCC and the 2010 data set, neutrality test statistics were calculated only for fragments with at least 300 analyzed nucleotide sites.

The distribution of polymorphism neutrality test statistic values for the 238 analyzed CCC fragments was compared with the genome-wide distribution of the same statistics estimated from 990 fragments from the 2010 data set. The graphs of the distributions are normal Kernel smoothing plots for the two-point average interval frequency scatter. We used the two-sample smooth test of Janic-Wróblewska and Ledwina (2000) to test the similarity between the two distributions. This statistic models a distribution in k dimensions and has been shown to be a powerful test to compare distributions (Ducharme and Ledwina, 2003). Moreover, if the test is significant, the k different components in the decomposition of the statistic give additional information about how the distributions differ. We analyzed the smooth plots to the order $k = 4$. The first component, which is exactly the Mann-Witney statistic, is related to a difference in location, the second component to difference in scale, the third to difference in skewness, and the fourth to difference in kurtosis.

For Tajima's D and Fu and Li's D and F, we identified outliers in the empirical distributions of the statistics. Under the assumption of neutrality for the vast bulk of the genome-wide fragments, the ranked distribution of neutrality test statistic values calculated for the data set of 2010 fragments can be considered as the null. The test statistic values at the negative and positive 2.5 percentiles of the genome-wide distribution are

set as thresholds. Neutrality test statistics values for the 238 analyzed CCC fragments were considered significant if they exceeded the threshold value, referred to as a rejection of neutrality at the 5% significance level. This nonparametric empirical approach is not a formal test of neutrality, and values in the 2.5% tails are interpreted as outliers in the empirical distribution, where we must accept a reasonable chance for false positives (Kelley et al., 2006). However, for *A. thaliana*, it is considered as the preferred approach to follow (Schmid et al., 2005; Tenaillon and Tiffin, 2008) and has been used in a number of studies (Carlson et al., 2005; Toomajian et al., 2006; Borevitz et al., 2007).

Accession Numbers

A complete list of accession numbers for the sequences in the CCC data set can be found in Supplemental Data Set 1 online.

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure 1. Correlation between θ_w and Gene Density.

Supplemental Figure 2. Visual Representation of *CYCB1;4*, *CDKD;3*, *CYCB1;2*, and *E2Fa* Haplotypes.

Supplemental Table 1. List of Resequenced *A. thaliana* Accessions.

Supplemental Table 2. Summary of Nucleotide Diversity.

Supplemental Table 3. Data for the CCC Frame Shifts.

Supplemental Table 4. Summary of Codon Site Statistics for the CCC Genes.

Supplemental Data Set 1. Summary of Sequence Data and Primer Sequences.

Supplemental Data Set 2. Summary of SNP Frequency Pattern Statistics for the CCC Genes.

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A Population Genomics Study of the Arabidopsis Core Cell Cycle Genes Shows the Signature of Natural Selection

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