



Functional and evolutionary genomic inferences in *Populus* through genome and population sequencing of American and European aspen

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The *Populus* genus is one of the major plant model systems, but genomic resources have thus far primarily been available for poplar species, and primarily *Populus trichocarpa* (Torr. & Gray), which was the first tree with a whole-genome assembly. To further advance evolutionary and functional genomic analyses in *Populus*, we produced genome assemblies and population genetics resources of two aspen species, *Populus tremula* L. and *Populus tremuloides* Michx. The two aspen species have distributions spanning the Northern Hemisphere, where they are keystone species supporting a wide variety of dependent communities and produce a diverse array of secondary metabolites. Our analyses show that the two aspens share a similar genome structure and a highly conserved gene content with *P. trichocarpa* but display substantially higher levels of heterozygosity. Based on population resequencing data, we observed widespread positive and negative selection acting on both coding and noncoding regions. Furthermore, patterns of genetic diversity and molecular evolution in aspen are influenced by a number of features, such as expression level, coexpression network connectivity, and regulatory variation. To maximize the community utility of these resources, we have integrated all presented data within the PopGenIE web resource (PopGenIE.org).

formation, ease of propagation in tissue culture, and rapid growth, *Populus* has been firmly established as an important model system for studies of forest tree species with a mature set of genetic and genomic resources (2, 3). Moreover, poplars are “replete with

Significance

We performed de novo, full-genome sequence analysis of two *Populus* species, North American quaking and Eurasian trembling aspen, that contain striking levels of genetic variation. Our results showed that positive and negative selection broadly affects patterns of genomic variation, but to varying degrees across coding and noncoding regions. The strength of selection and rates of sequence divergence were strongly related to differences in gene expression and coexpression network connectivity. These results highlight the importance of both positive and negative selection in shaping genome-wide levels of genetic variation in an obligately outcrossing, perennial plant. The resources we present establish aspens as a powerful study system enabling future studies for understanding the genomic determinants of adaptive evolution.

genome assembly | natural selection | coexpression | population genetics | *Populus*

The genus *Populus* comprises ~30 species, including poplars, cottonwoods, and aspens. *Populus* has a distribution spanning the Northern Hemisphere, and numerous species and hybrids have been extensively planted globally. Poplars and aspens are pioneer species with among the most rapid growth of any temperate tree species, partly as a result of their characteristic heterophyllous growth. These traits, which are enhanced in interspecific hybrids, render poplars of commercial value with end uses that include biofuel, fiber, timber, bioremediation, and animal feed (1). Poplars are readily amenable to vegetative propagation and, consequently, have been closely associated with agriculture since before the Middle Ages, having been used as windbreaks, to prevent soil erosion and to stabilize river banks. As a result of the relatively small genome (<500 Mbp), suitability for efficient genetic trans-

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Data deposition: All raw sequencing data have been deposited at the European Nucleotide Archive resource, <https://www.ebi.ac.uk/ena> (accession no. PRJEB23585), except for *Populus davidiana* sequencing data, which are available from the Populus Genome Integrative Explorer (PopGenIE.org) Lin2018 web resource (accession no. pnas201801437).

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genetic variation at many different levels: among sections within the genus, as well as among species, provenances, populations, individuals, and genes” (4) as a result of their obligate outcrossing (dioecious) nature and airborne dispersal of pollen and seed. These characteristics render *Populus* an ideal system for advancing our understanding of the transition from juvenile to mature and reproductive phases (5), the genetic architecture underlying natural variation of complex phenotypes, studies of local adaptation (6–9), and studies of the divergence continuum (10, 11). Such studies necessitate available genome assemblies and corresponding genomics and population genetics resources.

Here, we present de novo assemblies for two aspen species, *Populus tremula* and *Populus tremuloides*. In comparison to the high-quality assembly of the *Populus trichocarpa* genome, our study provides a starting point for comparative and evolutionary genomics in the field of forest trees. Despite many attributes that differentiate aspens and other poplars, we found that their ge-

nomes are remarkably conserved across species. However, the two aspen species display substantially higher levels of heterozygosity both within and between individuals compared with *P. trichocarpa*. Thus, in addition to examining gene and genome evolution in *Populus*, the genomes of these species provide an excellent model for studying how evolutionary processes affect patterns of genetic variation across genomes. Access to the dataset of these *Populus* genome sequences will not only facilitate studies concerning adaptive evolution in natural plant populations but also accelerate the pace at which the untapped reservoir of adaptive genes can be exploited to understand how widespread forest trees may respond to future climate change.

Results and Discussion

Genome Assembly and Annotation. Here, we report the results of the genome assembly, gene, and transposable element (TE) annotation of two aspen species, *P. tremula* and *P. tremuloides*

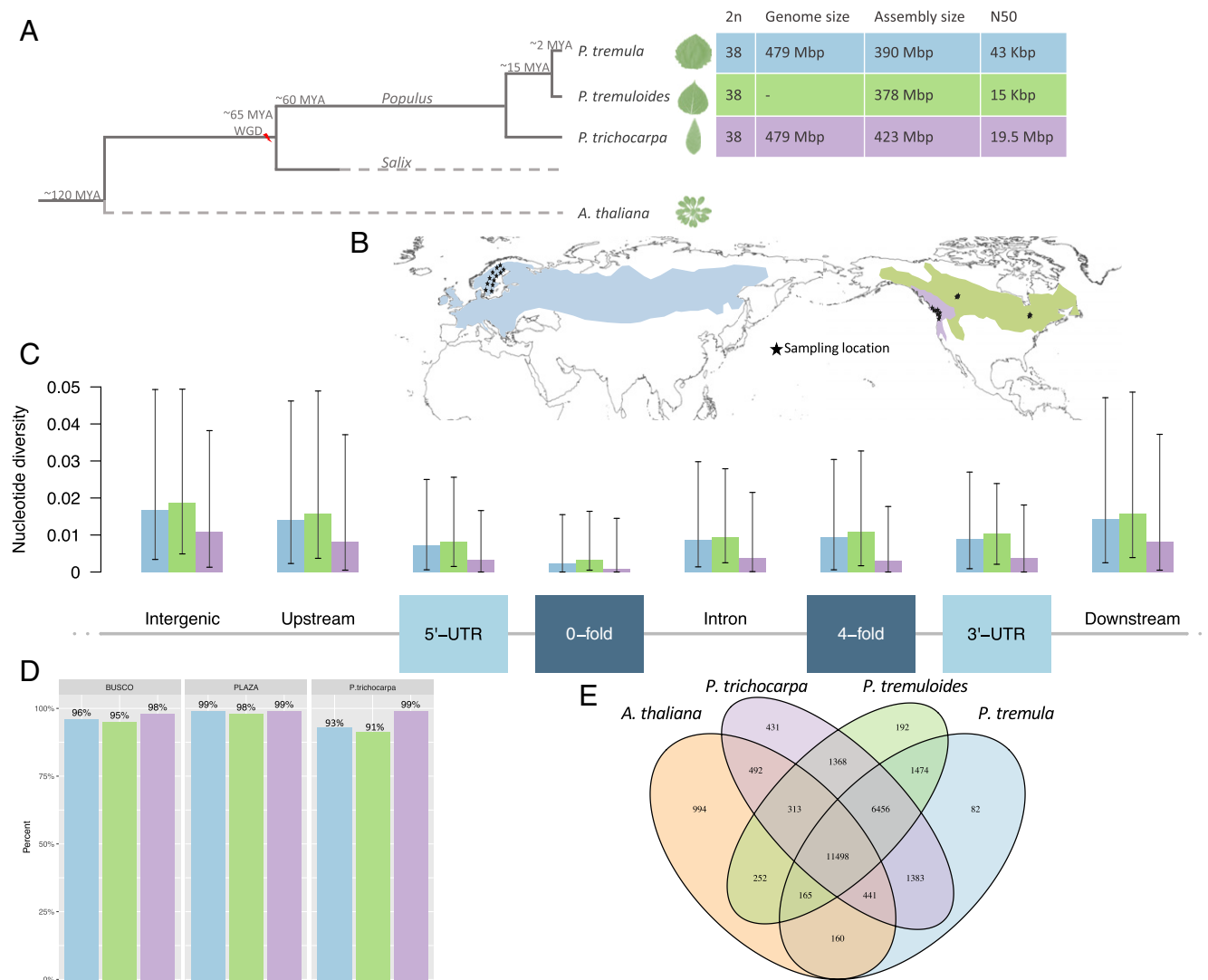


Fig. 1. Genome overview. (A) Simplified phylogram representing estimated divergence times (11, 13) and assembly statistics for the two genome assemblies presented here (*P. tremula* and *P. tremuloides*), with *P. trichocarpa* and *A. thaliana* indicated for reference. *P. trichocarpa* assembly statistics are based on Joint Genomes Institute genome release v3.0 of the Joint Genomes Institute. Both genomes were obtained from the Phytozome resource (<https://phytozome.jgi.doe.gov/pz/portal.html>). (B) Sampling localities (black stars) and distribution of *P. tremula*, *P. tremuloides*, and *P. trichocarpa*. (C) Nucleotide diversity in various genomic contexts calculated from alignments of resequencing data from 24 *P. tremula*, 22 *P. tremuloides*, and 24 *P. trichocarpa* individuals aligned to the corresponding genome assembly. (D) Percentage of genes represented in each *Populus* genome for BUSCO, PLAZA, and *P. trichocarpa* v3.0 gene sets. (E) Venn diagram representation of shared and unique gene families among *P. tremula*, *P. tremuloides*, *P. trichocarpa*, and *A. thaliana*.

(Fig. 1A and *SI Appendix* for all methods and details); comparative analyses of the two aspens to *P. trichocarpa*; and resequencing-based analyses of individuals from all three species (Fig. 1B). The *Populus* section, containing the aspens, diverged from the *Tacamachaca* section, containing the previously sequenced *P. trichocarpa* (12), ~15 Mya (13). Based on read alignments, we estimated that there is ~2% divergence between *P. tremula* and *P. trichocarpa* within coding regions (indicated by alignment mismatches). However, >50% of *P. tremula* reads could not be aligned to the *P. trichocarpa* genome, indicating extensive divergence between the two species within noncoding regions (*SI Appendix*, section 2.5.3). The aspen and *P. trichocarpa* genomes are of similar size, at about 480 Mbp (*SI Appendix*, Dataset S2), but a limiting factor for the aspen genome assemblies was the markedly higher nucleotide diversity, as revealed both by the k-mer spectra of unassembled reads and alignment of population resequencing reads (Fig. 1C and *SI Appendix*, Fig. S2.1). We assembled the *P. tremula* genome using a hybrid approach that merged 454 and two assemblies of Illumina short-read, paired-end (PE) libraries in a stepwise manner, the result of which was subsequently scaffolded using linking [mate-pair (MP)] libraries (*SI Appendix*, Fig. S2.3). The *P. tremuloides* genome was produced using a single assembly of Illumina short-read, PE libraries. Both assemblies contained >100,000 scaffolds and had relatively low contiguity (Table 1). Despite the aspen assemblies being relatively fragmented, 93% of *P. trichocarpa* (v3.0) genes were represented within the *P. tremula* assembly (*SI Appendix*, section 3.3.2), with the BUSCO (14) and PLAZA (15) gene sets further supporting high contiguity and completeness of the gene space in both aspen species (Fig. 1D and *SI Appendix*, section 3.3). Annotation of the gene space of *P. tremula* and *P. tremuloides* identified 35,984 and 36,830 genes, respectively (Table 1 and *SI Appendix*, section 3.2). Using ab initio and evidence-based methods to identify and annotate repetitive elements in the two aspen genomes (*SI Appendix*, section 3.1), we found that the majority of elements were long terminal repeat (LTR) TEs from the Ty1-Copia and Ty3-Gypsy families, in ad-

dition to a large number of unclassified TEs (Table 1). A phylogenetic analysis of the Ty1-Copia and Ty3-Gypsy families revealed no evidence for any species- or section-specific family members (*SI Appendix*, Figs. S3.1 and S3.2), demonstrating that the LTR complement predates the divergence of these species. Although long interspersed nuclear elements are rare in all of the genome, two clear examples of postspeciation amplification were identified for *P. trichocarpa* and, to a more limited extent, for *P. tremula* (*SI Appendix*, Fig. S3.3).

Comparative Genomics Identifies Extensive Conservation of the *Populus* Gene Space. On the basis of cross-species gene alignments, 37,238 *P. trichocarpa* genes were commonly aligned to both aspen species, whereas 1,127 genes were identified as putatively aspen-specific. In addition, 136 *P. tremula*, 146 *P. tremuloides*, and 536 *P. trichocarpa* genes were putatively species-specific (*SI Appendix*, section 4.2). A gene family clustering analysis placed >90% of genes into ~22,000 gene families, with 17,954 gene families shared among all three *Populus* species (Fig. 1E and *SI Appendix*, section 5.1). We identified 2,246 genes potentially under diversifying selection between aspen and *P. trichocarpa* [ratio of synonymous to non-synonymous number of nucleotide substitutions per site (K_a/K_s) > 1], which were enriched for gene ontology (GO) process categories, including “regulation of transcription,” “gene expression,” “biosynthetic process,” and “metabolic process” (*SI Appendix*, Dataset S1). Among these genes, there were a number of transcription factors from the NAC, MYB, YABBY, bHLH, and WRKY transcription factor families in addition to genes involved in cell wall biogenesis (xyloglucan endotransglucosylase/hydrolase and expansin), abiotic (LEA, heat shock proteins, and RD22) and biotic stress (LRR and disease resistance proteins), cell cycle and developmental regulators [cyclins, epidermal patterning factor, floral time, and homeotic control proteins (CONSTANS-like)], lipid transfer proteins, and an extensive number of proteins of unknown function. We examined the expression characteristics of genes with evidence of selection using population-wide RNA-sequencing data from the Swedish Aspen (SwAsp) collection (16) and the *P. tremula* tissue expression atlas (17). In the SwAsp gene coexpression network, genes potentially under diversifying selection on the basis of K_a/K_s between the two aspen species had slightly lower within-module (Mann–Whitney test, $P = 0.001$) and global (Mann–Whitney test, $P = 0.01$) network connectivity. This suggests that highly connected genes experience lower levels of diversifying selection than genes with fewer connections. The same was also true for genes potentially under diversifying selection on the basis of K_a/K_s from *P. trichocarpa*, but with a slightly more pronounced effect (Mann–Whitney $kdiff_norm$ $P = 0.0006$ and $kTotal$ $P = 0.03$; where $kTotal$ is the total connectivity for a gene within the network and $kdiff_norm$ is the difference between $kWithin$, which is connectivity of a gene within its assigned module, and $kOut$, which is the difference between $kTotal$ and $kWithin$, scaled for module size). This potentially reflects the greater divergence time between *P. trichocarpa* and *P. tremula*. Genes potentially under diversifying selection on the basis of K_a/K_s also had significantly higher absolute effect sizes for associated expression quantitative trait loci (eQTLs) (16), both for genes diverged between the two aspen species (Mann–Whitney $P < 2.2e-16$) and between *P. trichocarpa* and *P. tremula* (Mann–Whitney $P < 2.2e-16$). This is in agreement with the observation that eQTLs were enriched in genes with lower coexpression network connectivity at the periphery of the coexpression network (16). Together, these results indicate that genes with lower network connectivity are more likely to experience diversifying selection, and therefore to contribute to divergence among species driven by expression or regulatory modulation.

We performed comparisons of the genome sequences of the three species using an unsupervised method (18) that combines hidden Markov models and self-organizing maps to segment the

Table 1. Genome assembly, repeat, and gene space annotation summary statistics for *P. tremula* and *P. tremuloides*

Assembly	<i>P. tremula</i>	<i>P. tremuloides</i>
No. of scaffolds	216,318	164,504
Total size of scaffolds, bp	390,124,095	377,489,497
No. of scaffolds >500 bp	57,475	59,039
No. of scaffolds >1,000 bp	31,806	39,866
No. of scaffolds >10,000 bp	5,161	10,248
No. of scaffolds >100,000 bp	687	28
N50 bp	42,844	15,222
Repeat annotation (% values)		
Total	21.54	22.09
Ty1-copia	4	4.02
Ty3-gypsy	7.4	7.2
Other-LTR	0.13	0.11
LINEs	0.38	0.35
SINEs	0.36	0.38
DNA	3.28	3.43
NHF	5.99	6.6
Gene annotation (counts)		
High/low-confidence gene loci	29,252/6,057	26,842/8,852
High/low-confidence transcripts	76,557/8,312	34,439/13,899
High/low gene loci expressed	27,825/4,833	NA/NA

LINEs, long interspersed nuclear elements; N50, scaffold length for which at least half of the nucleotides in the assembly belong to scaffolds with the N50 length or longer; NA, not applicable; NHF, no hit found, i.e., elements that are LTR-RTs related but do not have significant similarity with the major families; SINEs, short interspersed nuclear element.

P. tremula (11), 22 *P. tremuloides* (11), and 24 *P. trichocarpa* (7) individuals (SI Appendix, section 7). We aligned reads from each individual to their respective assembled genome, which enabled us to access a greater proportion of the genomes after filtering (77.6% for *P. tremula* and 82.1% for *P. tremuloides*) than alignments to the *P. trichocarpa* assembly (42.8%; SI Appendix, section 7). We observed that both aspen species harbor substantially higher genome-wide levels of genetic diversity compared with *P. trichocarpa* (Fig. 1B and SI Appendix, Table S7.2). This can likely be ascribed to different demographic histories and larger distribution ranges that collectively lead to a higher effective population size (N_e s) in the two aspen species compared with *P. trichocarpa* (11, 24). The allele frequency spectrum of polymorphic sites also differed between species, with a more pronounced excess of low-frequency polymorphisms (negative Tajima's D value) in aspens than in *P. trichocarpa* (SI Appendix, Table S7.3). This is in accordance with analyses indicating that both aspen species have experienced recent range expansions following the last glaciation (25). Nucleotide diversity showed consistent patterns of variation across coding and noncoding regions in the three species, with diversity being approximately three- to fourfold higher at fourfold synonymous and noncoding sites than at zero-fold nonsynonymous sites (Fig. 1B and SI Appendix, Table S7.2). Furthermore, population-scaled recombination rates were inferred to be substantially higher in aspens, particularly in *P. tremuloides* (SI Appendix, Fig. S7.3), which had the lowest levels of LD (SI Appendix, Fig. S7.2), likely reflecting historical differences in effective population sizes between aspens and *P. trichocarpa*. Interestingly, genomic regions in close proximity to genes exhibited the highest recombination rates in all three species, (SI Appendix, Fig. S7.3). We also identified insertions and deletions (INDELs) using the same resequencing data (SI Appendix, section 8), finding that >50% of identified INDELs were common to *P. tremula* and *P. tremuloides*, suggesting that they occurred after the *Populus* section split from the *Tacamahaca* section but before the speciation event that separated *P. tremula* and *P. tremuloides* ~2.3 Mya (11). Short INDELs (<100 bp) were rare in coding regions (1–2%), with a higher proportion of long INDELs (>100 bp) affecting coding regions (15.2%), and, similar to SNPs,

there was an excess of low-frequency variants at INDELs, as indicated by negative Tajima's D values (SI Appendix, Fig. S8.3).

We further used the DFE-alpha approach (26, 27), which jointly infers demographic and selective parameters using polymorphism and divergence data, to quantify the impact of negative and positive selection on sites located in different genomic contexts in the two aspen species (SI Appendix, section 7.2). For both aspen species, we found that negative selection was substantially stronger in coding than noncoding regions, with more than 40% of zero-fold nonsynonymous sites being subject to strong negative selection ($N_e s > 100$; Fig. 3A and Table 2). In noncoding regions, 5' UTRs showed stronger negative selection than other regions (Fig. 3A), with ~30% of 5' UTRs under moderate negative selection ($1 < N_e s < 100$) in both aspen species. In addition, we found widespread evidence of positive selection in coding regions, where a high proportion of divergence at zero-fold sites has been driven to fixation ($\alpha = 30\text{--}40\%$), corroborating a previous study based on far fewer genes (6). Similarly, in both aspen species, there were high proportions of substitutions fixed due to positive selection ($\alpha = 30\text{--}40\%$) and higher rates of adaptive substitutions relative to neutral divergence in 5' UTRs ($\omega \sim 30\%$; Fig. 3A), mirroring results from *Drosophila melanogaster* (5) and *Capsella grandiflora* (28). Through analysis of orthologous promoter regions among the three species, we found that genes with highly conserved upstream and downstream regions were enriched for DNA binding activity (SI Appendix, section 4.3), suggesting that these regions are functionally important, and hence provide an explanation for why they are subject to both strong purifying selection and adaptive evolution (Fig. 3A). We caution that estimation of α could be biased if ancient demographic fluctuations are not sufficiently well captured by current polymorphism data (29).

Gene Expression Characteristics Influence Functional Constraint and Adaptive Evolution. As gene expression and regulation changes have been postulated to be key determinants of rates of adaptive evolution (30, 31), we tested for the relative contributions of negative and positive selection in driving differences in rates of

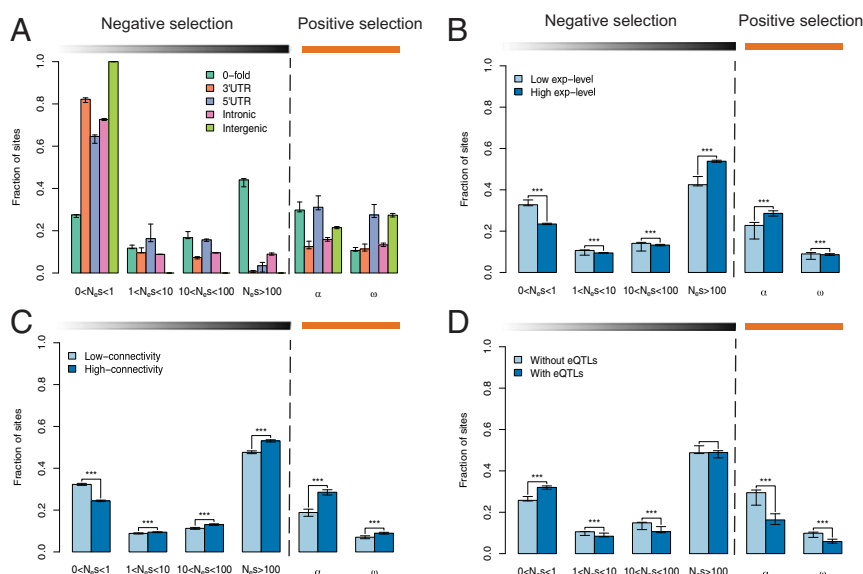


Fig. 3. Population genetics for *P. tremula*. (A) Estimates of negative and positive selection on coding and noncoding regions, separated by site type. Error bars represent 95% bootstrap confidence intervals. Estimates of negative and positive selection on zero-fold nonsynonymous sites in genes with varying expression level (B), varying connectivity level in coexpression network (C), and with or without eQTLs (D) are shown. $N_e s$ categories represent different bins of negative selection strength. α , proportion of divergent sites fixed by positive selection; ω , rate of adaptive substitution relative to neutral divergence. All calculations are based on genomic resequencing of 94 *P. tremula* individuals with reads aligned to the *P. tremula* genome assembly. *** $P < 0.001$.

suggests stronger negative selection acting on nonsynonymous sites in more highly expressed and highly connected genes (16, 34). Additionally, the strength of negative selection acting on these genes was even stronger after controlling for differences in selection on synonymous sites (*SI Appendix, section 7.3 and Dataset S2*). These patterns were also supported by the regression model, which showed that most expressed genes in the *P. tremula* genome have evolved under selective constraint (18,612 of 22,306 genes; Fig. 4A), and that genes evolving under selective constraints have significantly higher expression levels and contained a greater fraction of core genes [hubs in coexpression network modules (16)] than genes that are evolving neutrally (Fig. 4B). In comparison to the clear evidence observed at nonsynonymous sites, selection on noncoding sites was not consistent before and after controlling for the synonymous site selection, which could be due, in part, to the general weaker purifying selection on noncoding than coding regions (*SI Appendix, section 7.3 and Dataset S2*). We thus focused on nonsynonymous sites in the remainder of our analyses.

We found genes with high expression and high connectivity to have significantly higher proportions of adaptive nonsynonymous substitutions (α) than genes with low expression and low connectivity (Fig. 3B and C). However, we did not observe higher rates of adaptive substitutions (ω) in genes with high expression (Fig. 3B), and there was significantly lower expression among the 395 genes identified as being under positive selection by the regression analysis (Fig. 4C and D). Together, these results indicate that the higher α we observed for highly expressed genes most likely arises from an increased strength of negative selection acting on genes with high expression that eliminates a substantial fraction of weakly deleterious alleles rather than from an increased rate of fixation of adaptive mutations per se. In contrast to the pattern observed in expression level, genes with high connectivity show higher rates of ω compared with genes with low connectivity (Fig. 3C), and there was also a significantly higher proportion of core genes among the positively selected genes identified by the regression analysis (Fig. 4D). Although the positively selected genes from the regression analysis in *P. tremula* were enriched for the GO terms “signal transduction,” “cellular response to stimulus,” and “metabolic process” (*SI Appendix, Table S7.7*), we found that positively selected core genes were highly enriched in nonstructural carbohydrate metabolic and biosynthetic process, biotic stimulus response, and nucleosome and chromatin assembly (*SI Appendix, Table S7.8*), which may play critical roles in the resilience to stress and life history strategies of long-lived trees (35). Given that core genes are more likely to be required for network integrity and adaptive regulatory evolution than noncore genes (36), the characteristics of both stronger negative and positive selection on core genes may have promoted more rapid and efficient adaptation during the evolutionary history of long-lived forest trees such as aspen.

In addition to expression level and network connectivity, gene expression variation and associated regulatory loci are major contributors to phenotype variation in complex traits, and are thus likely to be subject to selection pressures (37–39). We found that coding sequence evolution in genes with high expression variance among genotypes tended to be under slightly weaker purifying selection and stronger positive selection compared with genes with low expression variance (Fig. 4B and D and *SI Appendix, Dataset S2*). Our previous study showed that there is pervasive regulatory variation in *P. tremula* (16), and we extend this observation here by finding that genes harboring regulatory variation (with identified eQTLs; eGene) have significantly higher proportions of nearly neutral nonsynonymous mutations (Fig. 3D and *SI Appendix, Dataset S2*), significantly lower proportions of adaptive nonsynonymous substitutions, and lower rates of positive selection than other genes (non-eGene; Fig. 3D). In agreement with a recent study in *C. grandiflora* (40), our

findings support the view that genes with regulatory variation, which we know are less central in the coexpression networks (16, 40), are evolving under weaker negative selection and undergo less adaptive evolution compared with genes without regulation variation. Taken together, our results show there is complex interaction between gene sequence and expression evolution in *Populus*, and future studies should explore the underlying driving evolutionary forces in greater detail across a wider range of plant species.

Conclusions

Here, and in other recent work, we have demonstrated the power of the aspen genome resources for understanding how natural selection varies among genes differing in gene expression (16) and how effects of linked selection vary across the genome within and among species (25) to help decipher the history of speciation in aspens (11) and identify the genomic basis of local adaptation (8). By developing an extensive genomic resource for aspen, our goal is to enable functional, comparative, and evolutionary genomic analyses in *Populus* and to extend the utility of the genus as a unique study system in plant and evolutionary biology. *Populus* has a number of features that makes it novel compared with many other model systems, such as an obligate outcrossing mating system, great longevity, abundant genomic diversity, and species of contrasting effective population size. To facilitate future studies of the many novel aspects of *Populus* biology, we have integrated all data generated here into the *Populus* Genome Integrative Explorer (PopGenIE.org) (17) web resource. These data will serve researchers performing evolutionary and comparative analyses as well as functional genomics studies aimed at deciphering genes underlying complex phenotypes through, for instance, aiding with the design of CRISPR guide-RNAs for gene editing.

Materials and Methods

Biological Materials. *P. tremula* genome sequencing was performed on DNA extracted from propagated root cuttings of a single wild tree growing on the Umeå University campus (63° 49' 17" N, 20° 18' 40" E). *P. tremuloides* DNA was extracted from mature, freeze-dried leaves from genotype Dan2-1B7. All DNA was extracted using a DNeasy Plant Mini Kit (Qiagen). Nuclear DNA contents were estimated by propidium iodide flow cytometry.

Sequencing. *P. tremula* DNA was sequenced as single-end reads using the Roche 454 platform and as PE and MP read libraries using Illumina short-read platforms. *P. tremuloides* DNA was sequenced using PE sequencing libraries on the Illumina HiSeq platform. *Populus davidiana* sequencing data were obtained from the National Institute of Forest Science, Korea. DNA was extracted from mature leaves sampled from mature trees in a common garden experiment in Suwon, Korea. We have deposited all raw sequencing data at the European Nucleotide Archive (ENA) resource (<https://www.ebi.ac.uk/ena>) as accession no. PRJEB23585, except for *P. davidiana* sequencing data, which are available from the PopGenIE.org web resource and will be described fully elsewhere.

Assembly. We used a hierarchical approach to assemble the *P. tremula* that comprised separate contig assemblies of the 454 and Illumina data that were merged in a stepwise manner, with the final merged assembly subsequently scaffolded using the MP libraries. As we only had PE Illumina data available for *P. tremuloides*, a single PE assembly was performed. We evaluated gene space coverage of the *P. tremula* final and substage assemblies by aligning the primary transcripts of the *P. trichocarpa* reference genome to the various assemblies. We used feature response curves (41) to assess assembly correctness.

Annotation. To annotate the repetitive fraction of the genome, we searched the *P. tremula* and *P. tremuloides* genome assemblies using RepeatScout (42) and retrieved *P. trichocarpa* repetitive sequences from Repbase (43), merging these to form a repeat library. We then used RepeatMasker (44) to identify repetitive elements in the genome assemblies. To annotate the protein-coding fraction of the genome, we performed gene space annotation in three steps: using MAKER (45) to generate gene annotations, which we iteratively refined using PASA (46) on the basis of four transcriptome

datasets before final manual curation. To assess gene space completeness, we used CEGMA (47), BUSCO (14), PLAZA coreGF (15), and alignments of the *P. trichocarpa* annotation. Cross-species gene alignments were performed using GMAP (48) to identify common and species-specific genes.

Comparative Genomics. To determine the orthologous relationship between the three *Populus* species, we collated protein sequences from *Arabidopsis thaliana* (TAIR10), *P. trichocarpa* (V3.0), and the two aspen species presented here, and performed an all-against-all BLASTp (49) sequence similarity search, the results of which were used to perform two rounds of clustering using TribeMCL (50) to delineate gene families. We performed multiple sequence alignments of each family and used these to calculate K_s and K_a , and their rate ratio (K_a/K_s). We aligned the aspen and *P. trichocarpa* genomes using Satsuma (51). To pseudoscaffold the aspen assemblies, we ordered and oriented the scaffolds from each assembly according to synteny to *P. trichocarpa* using Chromosome from the Satsuma package. We then generated syntenic self-alignments using Satsuma and visualized these using ChromosomePaint (52).

Population Genetics. We used whole-genome resequencing data from 24 genotypes of *P. tremula*, 22 genotypes of *P. tremuloides*, and 24 genotypes of *P. trichocarpa* to perform population genetics analyses. We used Trimmomatic (53) to remove adapter sequences and low-quality bases, after which all reads were mapped to their respective genome assembly using bwa-mem (54). SNPs were called using HaplotypeCaller from GATK (55). We used ANGSD (56) to estimate average pairwise nucleotide diversity (Θ_s) and Tajima's D for different genomic contexts (zerofold nonsynonymous, fourfold synonymous, intron, 3' UTR, 5' UTR, upstream and downstream regulatory regions, and intergenic sites) over nonoverlapping 1-Kbp windows. To estimate the rate of LD decay, we used PLINK (57) to randomly thin the SNPs and to calculate the squared correlation coefficients (r^2) between all pairs of SNPs within a distance of 20 Kbp. Using fourfold synonymous sites as the neutral reference, we estimated the distribution of fitness effects ($N_e s$), the proportion of fixations driven by positive selection (ω), and the rate of positive selection (ω) for each category of functional elements using DFE-alpha. To quantify the impact of negative and positive selection on different genomic regions, we used read alignments and SNP calling against the *P. trichocarpa* reference genome by

comparing the site frequency spectra and divergence of different genomic contexts with those for fourfold synonymous sites, which we assumed to be neutral. We binned all expressed genes (22,306 genes) from population-wide RNA-sequencing data in winter buds (16) into two subsets according to four different features related to gene expression to examine the relation of expression to signatures of selection. We ran an unsupervised genome-wide population analysis using Saguaro (18) to segment the genomes into different local topologies based on reads from all populations mapped to *P. trichocarpa*.

Structural Variants. We used a combination of Samtools (58) and Varscan (59) to detect short INDELS within mapped reads, Control-FREEC (60) to detect copy number variants based on sequence depth, Breakdancer (61) and Delly (62) to detect structural variants (SVs) based on altered distance between mapped reads from PE data, and Lumpy (63) to integrate multiple SV signals jointly across multiple samples. All methods were run with default parameters. The output from the different methods was combined using an in-house Perl script, and only variants detected by at least two methods were kept for further analyses. Variants were annotated using ANNOVAR (64).

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