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The first two chromosome-scale genome assemblies of American hazelnut enable comparative genomic analysis of the genus *Corylus*

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Summary

The native, perennial shrub American hazelnut (Corylus americana) is cultivated in the Midwestern United States for its significant ecological benefits, as well as its high-value nut crop. Implementation of modern breeding methods and quantitative genetic analyses of C. americana requires high-quality reference genomes, a resource that is currently lacking. We therefore developed the first chromosome-scale assemblies for this species using the accessions 'Rush' and 'Winkler'. Genomes were assembled using HiFi PacBio reads and Arima Hi-C data, and Oxford Nanopore reads and a high-density genetic map were used to perform error correction. N50 scores are 31.9 Mb and 35.3 Mb, with 90.2% and 97.1% of the total genome assembled into the 11 pseudomolecules, for 'Rush' and 'Winkler', respectively. Gene prediction was performed using custom RNAseg libraries and protein homology data. 'Rush' has a BUSCO score of 99.0 for its assembly and 99.0 for its annotation, while 'Winkler' had corresponding scores of 96.9 and 96.5, indicating high-quality assemblies. These two independent assemblies enable unbiased assessment of structural variation within C. americana, as well as patterns of syntenic relationships across the Corylus genus. Furthermore, we identified high-density SNP marker sets from genotyping-by-sequencing data using 1343 C. americana, C. avellana and C. americana × C. avellana hybrids, in order to assess population structure in natural and breeding populations. Finally, the transcriptomes of these assemblies, as well as several other recently published Corylus genomes, were utilized to perform phylogenetic analysis of sporophytic self-incompatibility (SSI) in hazelnut, providing evidence of unique molecular pathways governing self-incompatibility in Corylus.

Introduction

Species of hazelnut (Corylus spp.) are cultivated globally, and generate an annual production of 1.1 million tons of in-shell nuts, produced across 34 countries (FAOSTAT, 2023). This production is driven by a wide range of market uses across diverse industries, including food products, pharmaceuticals and dietary supplements (Ceylan et al., 2022). Currently, commercial cultivation relies overwhelmingly on cultivars of European hazelnut (C. avellana) adapted to Mediterranean climates (Di Lena et al., 2022) whose moderate winters satisfy their chilling requirements (Mehlenbacher, 1991). The distribution of hazelnut production reflects these climatic requirements, with major centres of production in Turkey, Italy and Spain, with Turkey alone producing over 65% of the global supply (Semih Uzundumlu et al., 2022). In the United States, this narrow range of adaptation currently restricts production to relatively small regions such as the Willamette Valley in Oregon (Revord et al., 2020).

As a woody perennial crop, hazelnuts have substantial potential to provide ecosystem services such as reduced soil erosion and nutrient runoff (Demchik *et al.*, 2014), while also sequestering carbon in above- and below-ground biomass

(Granata et al., 2020). Expanding the climatic conditions under which commercially viable hazelnut varieties can be grown would not only make production more resilient in the face of changing climates, but provide high-value alternative crops to producers across wider geographies. One possible method for the development of commercially viable hazelnut varieties adapted to novel growing regions is the improvement of the native North American shrub C. americana, which possesses disease resistance and cold hardiness traits lacking in most cultivated forms of European hazelnut (Molnar et al., 2018). Genomics-assisted breeding of C. americana oriented towards such improvement efforts, utilizing methods such as markerassisted selection and genomic prediction, would be greatly aided by a high-quality reference genome assembly for the species. Such a resource would, for example, facilitate the rapid identification and alignment of genetic polymorphisms across experimental and breeding populations (Kang et al., 2016). In addition, well-annotated assemblies provide the opportunity to investigate the genetic features underlying specific biomolecular pathways involved in abiotic and biotic stress resistance, enabling methods ranging from fine mapping of quantitative trait loci (QTL) to gene editing (Bolger et al., 2017; Dmitriev et al., 2022).

To facilitate such applied and experimental approaches, this study reports the first chromosome-scale reference assemblies for C. americana, built from two accessions - 'Rush' and 'Winkler' - which have historically been widely used in hazelnut breeding in the Eastern United States, largely as sources of resistance to the endemic fungal pathogen Anisogramma anomala (Eastern Filbert Blight; EFB) (Bhattarai et al., 2017). Constructed using a combination of long-read sequencing, chromosome conformation capture and genetic mapping, these chromosome-scale reference assemblies are of high guality and can immediately be deployed to advance breeding and genetic research objectives. The potential of these pseudohaploid assemblies is illustrated in this study through the identification of high-density single nucleotide polymorphism (SNP) markers for over 1343 hazelnut plants, drawn from Midwestern natural and breeding populations. These markers provide a detailed assessment of population structure in C. americana and interspecific hybrids between C. americana and C. avellana. 20th-century hazelnut breeding in the Midwestern United States involved interspecific hybridization with C. avellana (Rutter, 1987; Weschcke, 1954), but generations of open pollination and lack of detailed pedigree records obfuscate the genetic background of current Midwestern varieties. An analysis of population structure helps to resolve this question. Understanding the proportional representation of C. americana and C. avellana genetic backgrounds in varieties which have been successfully selected under Midwestern conditions will help determine breeding strategies for key traits.

In addition, the last 2 years have seen the release of chromosome-scale genome assemblies for several other *Corylus* species: *C. avellana* (cultivars 'Tombul' (Lucas *et al.*, 2021) and 'Tonda Gentile della Langhe' ('TGdL') (Pavese *et al.*, 2021)), *C. heterophylla* (accessions from Siping City, Jilin (Liu *et al.*, 2021)) and Yanqing, Beijing (Zhao *et al.*, 2021)) and a wild specimen of *C. mandshurica* (Li *et al.*, 2021). Together with the two *C. americana* genomes reported here, this study presents the first comparative genomic analyses across nearly half of the *Corylus* genus.

Comparative genetic analysis also is useful for analysing specific traits, such as self-incompatibility, an understanding of which is critical to both hazelnut breeding and production. Plants have evolved numerous strategies to increase the frequency of outcrossing, ranging from variation in the timing of floral development, to genetically regulated mechanisms. With respect to the latter, two primary classes of genetic self-incompatibility have been identified: gametophytic self-incompatibility (GSI) and sporophytic self-incompatibility (SSI) (Silva and Goring, 2001). The former is typified by Rosaceae and Solanaceae species, wherein the haploid genotype of the male gamete at an 'S-locus' (often encoding an F-box (SLF/SFB) protein) is detected by the female parent, which blocks the growth of the germinating pollen tube (Sassa, 2016; Sijacic et al., 2004). The latter is exemplified by Brassicaceae species, where the diploid genotype of the male parent at the S-locus (often encoding a cysteine-rich protein (SCR/SP11)) mediates detection by the female parent and a suppression of pollen germination (Hiscock and McInnis, 2003). Self-incompatibility in Corylus represents a form of SSI, regulated by a single S-locus (Mehlenbacher and Thompson, 1988). Fluorescent microscopic observation of pollen tube growth has identified 33 alleles with 8 levels of linear dominance at this S-locus (Mehlenbacher, 1997, 2014). We apply a phylogenetic approach to the analysis of SSI using the transcriptomes reported

here, and this analysis sheds additional light on the evolution of self-incompatibility in *Corylus*.

Results

Sequencing and assembly

Table 1 summarizes the raw sequence data used in assembling the two genomes. For 'Rush', 14.97 Gb of circular consensus sequencing (CCS) reads (44.32× coverage) and 13.60 Gb of ONT sequence (40.29× coverage) were generated. For 'Winkler', 19.60 Gb of CCS reads (58.02×) and 12.69 Gb of ONT sequence (37.57× coverage) was generated. Short-read sequencing of Arima Hi-C libraries generated $78.7 \times$ and $114 \times$ coverage for 'Rush' and 'Winkler', respectively. While 'Winkler' libraries, therefore, generated substantially more data across all sequencing platforms, in the case of both accessions, coverage was judged to be more than sufficient for the assembly of a diploid organism, assumed to have a relatively small genome size of ~350-370 Mb, based on comparisons with previous Corylus assemblies (Li et al., 2021; Liu et al., 2021; Lucas et al., 2021; Pavese et al., 2021; Zhao et al., 2021). The contact map for 'Winkler', showing contigs scaffolded into chromosome-scale pseudomolecules, is shown in Figure 1a (visualized using Juicebox (https://github.com/aidenlab/Juicebox)).

Table 2 presents statistics summarizing the quality and contiguity of the 'Rush' and 'Winkler' assemblies. While both genomes exhibit high quality and contiguity, with N50 scores >30 Mb, and over 90% of the genome assembled into 11 pseudomolecules, for both accessions there are clear differences which reflect the divergence in assembly methods. Both assemblies are close in length to previous reports of the size of Corylus species' genomes; however, the 'Rush' assembly is over 50 Mb larger than the 'Winkler' assembly. This is likely due to unresolved duplication, apparent also in Figure 1b, due to the fact that Hi-C sequence data was not be included in the assembly, which is also reflected in the many more contigs present in 'Rush' compared to 'Winkler'. While including Hi-C data for 'Winkler' led to a reduction in total contigs from 398 (following hifiasm and haplotig purge) to the 264 reported in Table 1, including Hi-C data for 'Rush' led to a fragmentation of the assembly, increasing the number of contigs to 1845. Benchmarking Universal Single Copy Orthologs (BUSCOs) were calculated for 'Rush' and 'Winkler': while both assemblies had nearly identical complete and single-copy BUSCOs represented (95.1% and 94.6%, respectively) 'Rush' contained more duplicates (3.8% vs. 1.4%).

This finding and the slightly larger maximum contig size also suggest unresolved artificial duplication. The duplicated regions visible in Figure 1B appear to be flanked by repetitive content, suggesting both haplotypes have been included in this draft

 Table 1
 Summary of sequence data generated on PacBio, Oxford

 Nanopore and Illumina sequencing platforms

Sample	Sequence type	Reads	Avg. length (bp)	Coverage
'Rush'	PacBio CCS	1 026 165	14 584	44.3×
'Winkler'	PacBio CCS	1 264 282	15 496	58.0×
'Rush'	ONT	581 538	22 053	40.3×
'Winkler'	ONT	616 903	21 814	37.6×
'Rush'	Arima Hi-C	88 601 323	2 × 150	78.7×
'Winkler'	Arima Hi-C	129 091 816	2 × 150	114.7×



Figure 1 (a) Contact map showing aligned and sorted Arima Hi-C data for 'Winkler' as visualized in Juicebox; green squares indicate ordered and oriented contigs, and blue squares indicate chromosomes. (b) Whole genome comparison of the 'Rush' and 'Winkler' chromosome-scale assemblies. Dotplots of raw windowed alignment were generated by GENESPACE. While small indels are evident, largely consistent synteny is observed across all 11 chromosomes. A single sizeable inversion on chromosome 9 is highlighted in red. Potential inclusion of both haplotypes is only evident in 'Rush'.

Table 2 Summary of statistics related to genome assemblies. BUSCOscore codes: C: complete; S: complete and single copy; D: completeand duplicated; F: fragmented; M: missing

Statistic	'Winkler'	'Rush'
Assembly size (bp)	337 645 099	388 211 906
Number of contigs	264	1118
GC content (%)	36.05	36.68
Assembly L50 (#)	5	5
Contig N50 (bp)	31 913 876	35 381 848
Maximum contig size (bp)	44 790 255	46 319 783
Genome in chromosomes (%)	97.1	90.2
BUSCO (C [S, D]; F; M) (%)	96 [94.6, 1.4]; 0.9; 3.1	98.9 [95.1, 3.8]; 0.2; 0.9

assembly. While not conclusive, these dotplots also highlight the fact that Winkler has no such identifiable duplications.

Both assemblies were annotated for several genomic features: gene models were predicted using a combination of RNAseq data and protein homology data; transposable elements were identified using RepeatModeler; tandem repeats were identified using TandemRepeatFinder; and tRNAs were predicted using tRNAscan-SE. These features are presented in a Circos-style plot in Supp. Figure S2. Specific statistics related to the gene model predictions are summarized in Table 3.

These annotations appear extremely similar in terms of the structure of the gene models that were predicted. Slightly fewer genes were predicted in 'Winkler', which in this case is likely not

Table 3 Summary of statistics for the annotation of 'Winkler' and 'Rush'

Statistic	'Winkler'	'Rush'	
Primary transcripts	23 331	24 562	
Alternative transcripts	5701	6174	
Median exon length (bp)	159	159	
Median intron length (bp)	245	248	
Average number of exons per gene	5.7	5.7	

the consequence of greater unresolved duplication in 'Rush', as the genomes were soft masked for repetitive elements. Finally, additional statistics related to the genome annotation are provided in Table 4, which compares the 'Rush' and 'Winkler' assemblies to the five currently published chromosome-scale *Corylus* assemblies.

The total number of predicted genes is within the range previously reported in *Corylus*. In addition, the BUSCO scores for 'Rush' and 'Winkler' are substantially higher than the scores reported for any previous *Corylus* genome annotation and are very close to the assembly BUSCO scores. This suggests not only that these annotations are highly complete, but also provides support for the accuracy of the predictive procedure described above.

Comparison across the Corylus genus

Comparisons across multiple species can be made simultaneously or in a pairwise manner. Using the five previously

Table 4 Comparison with annotations of previously published Corylus species. BUSCO scores refer to all complete representatives, both single and duplicated

Species and cultivar name or accession	Genome size (11 chromosomes)	Repeat content (%)	Predicted genes	Genome BUSCO	Annotation BUSCO
	· · ·				
C. americana ('Winkler')	327.7	48.4	23 331	96.9	96.5
C. americana ('Rush')	350.2	49.0	24 562	99.0	99.0
C. mandshurica	367.7	68.7	25 923	97.1	92.2
C. heterophylla (Beijing)	361.9	56.7	27 591	93.5	92.0
C. heterophylla (Jilin)	342.9	58.0	22 319	94.7	92.7
C. avellana ('TGdL')	373.1	41.5	27 791	96.2	92.0
C. avellana ('Tombul')	369.8	38.1	27 270	97.0	76.8

published Corylus assemblies, which contain comparable annotations to the genomes presented here, together with the outgroups silver birch (Betula pendula) and apple (Malus domestica), the GENESPACE package (Lovell et al., 2022) was utilized to visualize syntenic relationships in both manners, as well as construct a pangenome assembly anchored to C. americana 'Winkler'. Figure 2 illustrates the general result that across these Corylus genomes, there is extremely broad synteny between all defined blocks, illustrating the high degree of relatedness between each pair of species. Differences in pseudomolecule numbering in C. avellana 'Tonda Gentile della Langhe', C. heterophylla 'Jilin' and C. mandshurica is an artefact of chromosome numbering not being standardized across species. Instead, numbering is currently made according to descending physical contig size; while Corylus chromosome sizes are generally very similar, assemblies had slight variations in contig lengths. Figure S3 shows dot plots representing each pairwise comparisons between syntenic blocks for each of the seven Corvlus genomes. Despite the high degree of synteny across all Corylus species on a chromosome-scale, it is clear that C. americana exhibits nearly perfect synteny with the two C. avellana assemblies. This decreases marginally when compared against the two C. heterophylla assemblies, with small inversions evident on a number of chromosomes. C. mandshurica very clearly exhibits the least amount of synteny, with large inversions and rearrangements of syntenic blocks across most chromosomes.

With respect to *B. pendula*, this figure makes clear the specific large-scale rearrangements which relate the 14 chromosomes of *B. pendula* to the 11 of *Corylus* spp. Specifically, sections of chromosomes 1 and 3 in *B. pendula* are rearranged in chromosomes 1 and 4 in *Corylus* spp., with *B. pendula* chromosome 11 also being fused with the end of chromosome 1 in *Corylus*. In addition, chromosomes 4 and 6 in *B. pendula* together constitute chromosome 2 in *Corylus*, while chromosomes 12 and 14 constitute chromosome 3 in *Corylus*.

Single-copy orthologues unique to C. americana

It is also possible to use the orthogroups generated by OrthoFinder to determine, instead of synteny, specifically those putative genes in *C. americana* which possess no orthologue in any currently annotated *Corylus* species, nor the other two outgroups included in this analysis: *B. pendula* and *M. domestica*. This subset of the transcriptome consists of 66 predicted single-copy orthologues present in both 'Rush' and 'Winkler', but absent in all other analysed genomes (File S1). By filtering for

only those single-copy orthologues present in the predicted gene sets for 'Rush' and 'Winkler' limits the possibility that these unique genes are an artefact of spurious assembly errors in either of the independent 'Rush' or 'Winkler' assemblies. Similarly, including a diversity of assemblies of other *Corylus* species, as well as the *Betula* and *Malus* outgroups, limits the potential that these genes have been identified as unique simply as a result of more successful gene prediction for 'Rush' and 'Winkler', which due to the high BUSCO scores for the annotations reported here, would otherwise be a concern.

Many of these genes logically did not return hits when the Viridiplantae database was queried using NCBI-BLAST. Several, however, are predicted to be involved in defence response pathways. Of particular note is the gene CamerWink-ler.08G009300 (homologous to CamerRush.08G012000.1 in 'Rush'), which is characterized as involved in 'defense response to fungus'. Given *C. americana*'s high level of resistance to the endemic fungal pathogen EFB, this gene would be a valuable target for future functional characterization.

Contributions of C. *americana* and C. *avellana* to current Midwest germplasm

A total of 1343 individual plants were sequenced using GBS, and SNP markers were called using the TASSEL GBSv2 pipeline (Bradbury, 2007). Sampled plants represented wild C. americana, cultivars from breeding programmes in Oregon, New Jersey and Minnesota, and F₁ populations comprising controlled crosses between these cultivars. A biplot for the first two principal components of the distance matrix constructed using these markers is shown in Figure 3. This visualization makes evident the wide genetic diversity represented in current Midwestern hazelnut varieties and lends support to the significant C. avellana contribution to specific selections, such as Rose9-2. Many currently named Midwestern varieties, on the other hand, cluster extremely closely with 'Rush', 'Winkler' and wild C. americana from the DNR, suggesting that while an interspecific cross may have occurred in the past, subsequent selection of progeny greatly favoured C. americana genetic contributions.

Phylogenetic analysis of sporophytic self-incompatibility

The locus regulating sporophytic self-incompatibility (SSI) in hazelnut has been fine-mapped in *C. avellana* (Hill *et al.*, 2021) and *C. heterophylla* x *C. avellana* interspecific hybrids (Hou *et al.*, 2022). These studies identified three MIK2 homologues believed to be responsible for SSI in these two *Corylus* species. In



Chromosomes scaled by gene rank order

Figure 2 Riparian plot generated by GENESPACE. Single-copy orthologues were used to generate and visually compare syntemy between 'Rush' and 'Winkler', the five other published *Corylus* genomes, and the outgroups *Betula pendula* and *Malus domestica*. In addition to illustrating consistent genomewide inter-species syntemy across the 11 chromosomes of *Corylus*, this plot clearly illustrates the large-scale relationships between these 11 chromosomes and the 14 chromosomes of *B. pendula*.



Figure 3 PCA biplot of SNPs identified in *C. americana*, *C. avellana* and interspecific hybrids. On the left, wild *C. americana* sourced from the DNR clusters together with 'Rush' and 'Winkler', along with a number of Midwestern varieties and an F_1 family produced through a cross between two of them (light and dark green dots, respectively). In the middle, an outlier Midwest variety (Rose9-2) exhibits PC1 scores similar to an F_1 between Eric4-21 (a Midwest variety that appears similar to *C. americana*) x Jefferson (a *C. avellana* variety from Oregon State University). On the right are *C. avellana* varieties from Oregon State University and Rutgers University, along with an interspecific F_1 population between a Rutgers cultivar and Rose9-2. This plot illustrates that while most Midwestern varieties appear to be genetically quite similar to *C. americana*, there is substantial diversity among them (more so than what is observed between the limited number of *C. avellana* varieties included in the analysis) with some appearing quite similar to interspecific F_1 hybrids – a finding supported by the historical record of interspecific hybridization in Midwestern breeding efforts.

order to examine their phylogenetic relationship, multiple sequence alignment was performed between the top BLAST hit for each of these three genes and all currently available *Corylus* transcriptomes. *B. pendula* and *B. oleracea*, which both also exhibit SSI (Hynynen *et al.*, 2010; Kitashiba and Nasrallah, 2014), as well as *M. domestica*, a more distantly related species that is a well-studied example of gametophytic self-incompatibility (Cheng *et al.*, 2006; Minamikawa *et al.*, 2010), were included as outgroups. This phylogenetic tree is shown in Figure 4 and illustrates the genus-specific evolution of SSI in *Corylus*, represented by several clades containing orthologous sequences only found in *Corylus* species.

Discussion

These assemblies represent a critical step towards developing the use of next-generation sequencing data in the improvement of *C. americana*, while also enabling comparisons of structural variation across *Corylus* species.

Variable contiguity of two de novo assemblies

The fragmentation of the 'Rush' assembly by the 3D-DNA pipeline (using Arima Hi-C sequence data) was unexpected, given the widespread and successful use of this method in building chromosome-scale assemblies (Ghurye *et al.*, 2019), and in

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Figure 4 A phylogenetic tree showing the evolution of genes comprising the self-incompatibility locus in *Corylus* across the seven available *Corylus* genomes, with *B. pendula*, *M. domestica*, and *B. oleracea* included as outgroups. Node labels represent bootstrap values based on 100 permutations. Colour overlays illustrate evident clades, and highlight the fact that several lineages of S-locus genes are entirely specific to *Corylus*.

particular, the substantial improvements in contiguity that was observed when using this data to refine the initial 'Winkler' assembly. Two possible explanations for this discrepancy suggest themselves. First, the 'Rush' accession generated less absolute sequence compared to the 'Winkler' assembly on all sequencing platforms, and as a result, there was less sequence data to include in each bioinformatic step. The raw assemblies generated by hifiasm were as a result also more contiguous for Winkler compared to 'Rush'.

Nevertheless, with coverage >40× for all sequence data types, the failure of Hi-C sequence to improve contiguity would not appear to be solely a consequence of a lack of data. Indeed, when reads were randomly removed from the Hi-C sequence data for 'Winkler', to simulate the coverage of Hi-C reads obtained for 'Rush' (78.7×), no decrease in the contiguity of the resulting assembly was observed, relative to the assembly generated using

the full set of 'Winkler' Hi-C reads. Artificially reducing coverage of the 'Winkler' Hi-C sequence data to even $30 \times$ still resulted in an assembly with a larger N50, and equivalent L50, relative to the assembly prior to running 3D-DNA. The fragmentation of the 'Rush' assembly caused by the 3D-DNA pipeline, therefore, cannot solely be explained by the number of Hi-C reads generated. A second likely factor is a difference in the relative heterozygosity of 'Rush' and 'Winkler'. Heterozygous single-individual assemblies are more challenging to construct, relative to fully homozygous individuals (Garg, 2021). Using the 53 696 GBS-derived SNPs described above it was possible to directly estimate genome-wide heterozygosity as a ratio of heterozygous sites to total polymorphic sites. By this metric, 22% heterozygosity was observed for 'Rush', while only 17% heterozygosity was observed for 'Winkler', suggesting that this could have contributed to the challenges in scaffolding 'Rush' using Hi-C data.

Regardless of the specific underlying factors complicating this stage of genome assembly, it is clear that even in diploid species with relatively small genome sizes, integrating chromosomal conformation capture sequence data into genome assembly pipelines does not always lead to more contiguous assemblies, with the outcome likely dependent on sequence coverage, heterozygosity and the quality of the initial draft assembly. At the same time, the use of high-coverage long-read sequence data, together with modern assemblers for scaffolding and polishing chromosome-scale genomes, can generate high-quality reference assemblies, even in outcrossing, heterozygous organisms.

PCA-modelled population structure

This study reports the first combined analysis of relatedness between commercial varieties of hazelnuts from the three main breeding programmes in the United States: Oregon State University, Rutgers University and the Upper Midwest Hazelnut Development Initiative. This evaluation of population structure, by combining a diversity of clonal varieties with F₁ full-sib families produced through multiple pairwise controlled crosses of these clones, help elucidate a long-outstanding guestion regarding the degree to which Midwest-bred hazelnut accessions remain closely related to C. avellana. 20th-century Midwest breeding populations included interspecific crosses with C. avellana, but the precise frequency of such hybridization events, and the fact that they were often followed by several generations of open pollination within predominantly C. americana breeding orchards, has left this guestion unresolved. Figure 3 shows clearly that the majority of these Midwest accessions are indeed closely related to wild C. americana sourced from the Department of Natural Resources, as well as 'Rush' and 'Winkler' themselves. A clear outlier in this respect is the variety Rose9-2, which very closely resembles other F₁ interspecific crosses.

Cross-genus level analysis

Transcriptomic analysis of the seven Corylus genomes included in this study reveals a consistent distribution of genes across the 11 chromosomes. This analysis considers nearly half of the recognized species in Corylus (Thompson et al., 1996). The observed consistency in gene distribution suggests a relatively recent common ancestor of these hazelnut species. This finding is important for understanding the evolutionary relationships between these species and could have implications for hazelnut breeding and genetic improvement programmes by providing a foundation for identifying useful traits and genes that are present in all Corylus species. These observations should not, however, be considered a substitute for phylogenetic analyses, constructed either with genome-wide polymorphisms or organellar sequence data. Two such phylogenies have recently been reported (Helmstetter et al., 2019; Zhao et al., 2020), which suggest a closer relationship between C. americana and the Asian taxa, relative to C. avellana, which is a finding similar to that seen in the phylogenetic trees built using SSI genes presented above. These genome-wide syntenic relationships are not direct evidence against this finding, given that the algorithms for estimating phylogenetic relationships are not based on chromosome-scale evaluation of syntenic relationships in gene features. In addition, published phylogenies have included a far greater number of samples, relative to the limited number of chromosome-scale genome assemblies available for the comparative analyses carried out in this study. However, our results do provide important context when interpreting such phylogenies, demonstrating both a substantial level of genome-wide similarity across numerous clades within *Corylus*, as well as marginally greater synteny in the nuclear genome between species that are phylogenetically more diverged.

In addition to identifying general synteny on a genome-wide level, these analyses also made possible the identification of those single-copy orthologues with no representation in any other *Corylus* species. In particular, a gene found in both 'Rush' (CamerRush.08G012000.1) and 'Winkler' (CamerWinkler.08G009300) was ontologically associated with defence responses to fungal infection.

In particular, it appears to be a defensin-like (DEFL) gene, cysteine-rich antimicrobial proteins (Tesfaye *et al.*, 2013). This gene (CamerWinkler.08G009300) is, therefore, a promising target for future functional characterization. Quantitative genetic analyses have revealed numerous loci in *C. americana* that are associated with EFB resistance (Komaei Koma *et al.*, 2021; Lombardoni *et al.*, 2022; Revord *et al.*, 2020), and the gene reported here could be productively included in any future studies aimed at using expression analysis or forward genetic approaches to validating their role in plant defence pathways.

Sporophytic self-incompatibility

Attempts at genetically characterizing the SSI S-locus in Corylus have included hybridization-based staining of genes identified in Brassica oleracea L., which showed irregular hybridization (Hampson et al., 1996). Similarly, gene expression studies identified only 61% homology between B. oleracea alleles and S alleles in Corylus (Torello Marinoni et al., 2009). Finally, transcriptomic analysis of the fine-mapped S-locus region in multiple Corylus species suggest Corylus may harbour a novel SSI molecular mechanism that differs from Brassica (Hill et al., 2021; Hou et al., 2022). The analysis presented here suggests independent evolution of at least part of the sporophytic self-incompatibility mechanism that is unique to Corvlus and not shared by other members of the Betulaceae, nor model species of SSI such as *Brassica oleracea*. Only two implicated SSI genes, both from C. avellana 'Tombul', appear within a clade shared by B. oleracea, although occupying a relatively distinct sub-lineage. This shared ancestry, albeit limited, could suggest that a shared SSI system evolved prior to the species' divergence.

While gametophytic self-incompatibility (GSI) necessitates codominance between S alleles to prevent self-fertilization, SSI does not, allowing for the evolution of complex hierarchies of dominance and codominance (Brennan et al., 2011), characteristics observed in both Corvlus and Brassica (Mehlenbacher, 2014; Prigoda et al., 2005). Dominance in SSI has been implicated in accelerating rates of evolution within these systems, due to selective pressure for heterozygous genotypes to maximize their compatibility with the greatest number of Slocus genotypes (Schoen and Busch, 2009). This rapid rate of diversification within SSI systems may explain the inconclusive results observed in the gene tree in Figure 4 and makes the investigation of whether SSI evolved independently in the Brassicales and Fagales challenging. As long-read sequencing increasingly makes the assembly of high-quality, annotated genome assemblies more accessible to previously understudied species, future phylogenetic studies will hopefully be able to repeat the analyses performed here with a greater degree of resolution, helping to elucidate the specific point or points at which SSI systems evolved and diverged.

Conclusion

High-quality, annotated, chromosome-scale genome assemblies are essential tools for using modern genetic methods in the investigation of the molecular control of important traits, as well as the application of advanced breeding methods. These two assemblies provide an important resource which we hope will enable the application of such methods to the study and improvement of both *C. americana*, as well as interspecific hybrids which expand the range over which hazelnuts can be cultivated. Our analysis demonstrates highly conserved genomewide synteny across *Corylus* species. As such, genomic tools, analyses and resources developed in one species may be more broadly useful to breeding programmes in other species within the genus.

Experimental procedures

Plant material collection

Tissue was collected from C. americana accessions 'Rush' and 'Winkler' maintained at the National Clonal Plant Germplasm Repository (NCGR) in Corvallis, Oregon (PI 557022 and PI 557019, respectively). 'Rush' is a specimen collected around 1900 by J.F. Jones in Lancaster, Pennsylvania, while 'Winkler' was collected in 1910 by Wendell Williams in Danville, Iowa (Molnar, 2011). These two varieties were historically significant in early- and mid-20th century hazelnut breeding programmes and as such represent genetically relevant, pure C. americana selections, which also reflect the wide geographic distribution of C. americana in the Eastern United States. Today, 'Rush' remains a widely used source of EFB resistance (Bhattarai et al., 2017). Photos of the clones located at the NCGR are shown in Figure S1. Tissue was collected on April 13th, 2020, from bushes that had been etiolated for 3 days prior to sampling in order to minimize the concentration of volatiles and secondary metabolites in leaf tissue. Samples were immediately flash-frozen using liquid nitrogen and stored at -80 °C until DNA extraction.

Long-read sequencing

High-molecular-weight DNA was extracted following the protocol described by Vaillancourt and Buell (2019). In brief, leaf tissue was homogenized in liquid nitrogen, lysed with Carlson lysis buffer and purified with chloroform and Qiagen Genomic-tips (Qiagen N.V., Venlo, The Netherlands). Purity of extracted DNA was assessed spectrophotometrically using a NanoDrop[™] One (Thermo Fisher Scientific, Waltham, Massachusetts). DNA was quantified a using Qubit[™] dsDNA High Sensitivity kit (Thermo Fisher Scientific, Waltham, Massachusetts), and diluted and assessed for size using an Agilent FemtoPulse System (Santa Clara, California).

Pacific Biosciences HiFi libraries were prepared according to PN 101-853-100 Version 03 (Pacific Biosciences, Menlo Park, California). Modifications included shearing with a Covaris g-TUBE (Covaris, Woburn, Massachusetts) and size selecting with BluePippin (Sage Science, Beverly, MA). Libraries were sequenced on a PacBio Sequel II using the Sequel Polymerase Binding Kit 2.2 at the University of Wisconsin-Madison Biotechnology Center's DNA Sequencing Facility. Oxford Nanopore libraries were prepared following the Native Barcoding Expansion protocol (Oxford Nanopore Technologies, Oxford, United Kingdom). The library was sequenced on a R9.4.1 flowcell on an Oxford Nanopore PromethION, also at the UW-Madison Biotechnology

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Center. Partway through the sequencing run, DNA was flushed with the Oxford Nanopore Technologies Flow Cell Wash Kit (EXP-WSH004), and the additional library was loaded.

Hi-C sequencing

Nuclei were extracted from flash-frozen leaf tissue using a Sigma CelLytic[™] PN Plant Nuclei Isolation/Extraction Kit (Sigma-Aldrich, Burlington, Massachusetts). Crosslinking was performed following low-input protocols from Arima Genomics (Arima Genomics, Carlsbad, California). Crosslinked nuclei were quantified using a Qubit[™] dsDNA High Sensitivity kit, samples were sheared to 600 bp, and library preparation was performed using a KAPA® Hyper Prep kit (Roche, Basel, Switzerland). The final library was assessed for quality using the Agilent TapeStation System with a D1000 kit (Agilent Technologies, Santa Clara, California), and paired-end, 150 bp paired-end sequences were generated using an Illumina NovaSeq 6000 (Illumina, San Diego, California).

RNA extraction, library preparation and sequencing

Prior to RNA extraction, Tissuelyser II adapters, safelock tubes and 5 mm stainless steel beads were chilled at -80 °C for 2 h. Working on dry ice, leaf and kernel tissue was added to prechilled tubes and disrupted using the Tissuelyser II (Qiagen N.V., Venlo, The Netherlands) for 2 rounds at 30 Hz for 1 min each round. Samples were then processed using the Qiagen Plant RNeasy (Qiagen N.V., Venlo, The Netherlands) with an on-column DNA digest. Total RNA was assayed for purity and integrity using a NanoDrop One[™] Spectrophotometer and Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, California), respectively. RNA libraries were prepared from samples that met the Illumina TruSeg® Stranded Total RNA Sample Preparation Guide (15 031 048 E) input guidelines using the Illumina TruSeg® Stranded Total (Plant) RNA Sample Preparation kit (Illumina Inc., San Diego, California). For each library preparation, cytoplasmic, mitochondrial and chloroplast ribosomal RNA was removed using biotinylated target-specific oligos combined with paramagnetic beads tagged with streptavidin. Following purification, the reduced RNA was fragmented using divalent cations under elevated temperature. Fragmented RNA was copied into firststranded cDNA using SuperScript II Reverse Transcriptase (Invitrogen, Carlsbad, California) and random primers. Secondstrand cDNA was synthesized using a modified dNTP mix (dTTP replaced with dUTP). DNA Polymerase I and RNase H. Doublestranded cDNA was cleaned with AMPure XP Beads $(1\times)$ (Beckman Coulter, Brea, California). The cDNA products were incubated with Klenow DNA Polymerase to add a single 'A' nucleotide to the 3' end of the blunt DNA fragments. Unique dual indexes (UDI) were ligated to the DNA fragments and cleaned with two rounds of AMPure XP beads (0.8×). Adapter-ligated DNA was amplified by PCR and cleaned with AMPure XP beads $(0.8\times)$. Final libraries were assessed for size and quantity using an Agilent DNA1000 chip and a Qubit® dsDNA HS Assay Kit (Invitrogen, Carlsbad, California), respectively. Libraries were standardized to 2 nm, and 150-bp paired-end sequencing was performed on an Illumina NovaSeq 6000.

Genome assembly

An initial assembly was created using PacBio CCS reads with the program hifiasm v0.16.0-r369 (Cheng *et al.*, 2021) on a Unix server with 160 cores and 3TB of RAM. This draft assembly exhibited higher contiguity than references generated with Canu using ONT reads, HiCanu and Flye using either HiFi or ONT reads.

The program purge_haplotigs v1.1.1 (Roach *et al.*, 2018) was then utilized to reduce artificial genome duplication caused by incorporation of non-collapsed haplotigs in the final assembly. Hi-C reads were aligned, filtered and binned using the haplotigpurged assembly using the programme juicer (Durand *et al.*, 2016). Contigs were then scaffolded and ordered using the program 3D-DNA (Dudchenko *et al.*, 2017).

While integration of Hi-C data led to improved contiguity and reduced artificial duplication in the 'Winkler' assembly, contigs in 'Rush' were fragmented following the use of juicer and 3D-DNA. As a result, this step was omitted for 'Rush'. The assemblies were next iteratively polished three times with Racon (Vaser et al., 2017), using the original PacBio reads. As hifiasm performs polishing internally, this step was superfluous and to ensure it did not introduce artificial errors by re-inserting alleles from the alternate haplotype, we used mergury (Rhie et al., 2020) to confirm that the quality and completeness of the assembly were unaffected by Racon. We then used a recently constructed linkage map to detect erroneous inversions in the physical assemblies (Brainard et al., 2023), using visual inspection of heatmaps of recombination frequencies in an F₁ population. The program Ragtag v2.1.0 (Alonge et al., 2022), which is used to carry out reference-guided assembly, was then used to correct observed errors using ONT reads that overlapped any identified inversions. Initial guality assessments were made using BUSCO v5.4.3 (Manni et al., 2021) with the eudicots_odb10 database and guast v5.2.0 (Mikheenko et al., 2018) with default parameters.

Genome annotation

Transcript assemblies for both 'Rush' and 'Winkler' were made from ~200 M 2 × 150 bp stranded paired-end Illumina RNAseq reads using the software PERTRAN, which conducts genomeguided transcriptome short-read assembly via GSNAP (Wu and Nacu, 2010) and builds splice alignment graphs after alignment validation, realignment and correction. These outputs were subsequently used to construct 37 794 and 36 419 transcript assemblies for 'Rush' and 'Winkler', respectively, using the program PASA (Haas et al., 2003). Loci were determined by transcript assembly alignments and/or EXONERATE alignments of proteins from Mimulus guttatus, Arabidopsis thaliana, Gossypium raimondii, Betula platyphylla, Carya illinoinensis, Castanea dentata, Quercus rubra, Prunus persica, Fragaria vesca, Glycine max, Medicago truncatula, Vitis vinifera, Liriodendron tulipifera, Juglans microcarpa, Populus trichocarpa, Beta vulgaris, Solanum lycopersicum, Sorghum bicolor, Oryza sativa and Swiss-Prot (release 2022 04 of eukaryote proteomes), with up to 2 k bp extension on both ends unless extending into another locus on the same strand. Alignments were made to the respective genome assemblies following softmasking for repetitive elements. The repeat library consisted of de novo repeats identified by RepeatModeler2 (Flynn et al., 2020), generated using both the 'Rush' and 'Winkler' assemblies, as well as C. americana repeats identified in the databases RepBase and Dfam. Gene models were predicted by homology-based predictors: FGENESH+ (Salamov and Solovyev, 2000), FGENESH_EST (which is similar to FGE-NESH+, but uses EST to compute splice site and intron input instead of protein/translated ORF), EXONERATE (Slater and Birney, 2005), PASA assembly ORFs (a homology constrained ORF finder) and AUGUSTUS (Stanke et al., 2006) trained by the high confidence PASA assembly ORFs and with intron hints from short read alignments. The best-scored predictions for each locus were selected using multiple positive factors including EST and protein support and overlap with repeats as a negative factor. The selected gene predictions were improved by PASA. The improvement included adding UTRs, splicing correction and adding alternative transcripts. PASA-improved gene model proteins were subject to protein homology analysis to the above-mentioned proteomes to obtain a Cscore and protein coverage. Cscore is a protein BLASTP score ratio to the mutual best hit (MBH) BLASTP score, and protein coverage is the highest percentage of protein aligned to the best homologues. PASA-improved transcripts were selected based on Cscore, protein coverage, EST coverage and their CDS overlap with repeats. The transcripts were selected if their Cscore and protein coverage were ≥ 0.5 or if covered by ESTs. For gene models whose CDS were overlapped by repeats by more than 20%, their Cscore was required to be at least 0.9, and homology coverage at least 70%, in order to be selected. The selected gene models were subject to Pfam analysis, and gene models whose proteins were more than 30% overlapped by Pfam TE domains were removed as weak gene models. Incomplete gene models, low homology-supported gene models without full transcriptome support, short single exon (<300 bp CDS) with neither protein domains nor good expression, and repetitive gene models without strong homology support were manually filtered out.

Functional annotation

Every peptide sequence in the dataset was analysed with a computational pipeline that includes the standard InterProScan (Jones *et al.*, 2014) suite of programs to determine protein domains and other sequence features, E2P2 (Chae *et al.*, 2014; Schläpfer *et al.*, 2017) for enzyme assignments (EC) and PathoLogic (Karp *et al.*, 2021) for metabolic pathway assignments. Additional processing was used to determine Eukaryotic Orthologous Groups (KOG) gene assignment using a modified mutual best hit algorithm. Results of the InterProScan calculations were used to assign standard InterPro protein domain associations and from these, gene ontology (GO) terms. Protein domains inferred from these calculations were used to develop a putative gene functional assignment which includes a count of the multiplicity of the assignment in the proteome set.

Genotyping by sequencing

To better compare genomes of C. avellana and C. americana, we used genotyping-by-sequencing (GBS) to genotype a population of interspecific hybrids and a population of *C. americana*. Tissue from 1343 samples of breeding lines from the Upper Midwest Hazelnut Development Initiative (UMHDI), Oregon State University, in Corvallis, Oregon and Rutgers University, in New Brunswick, New Jersey, along with full-sib F₁ populations derived from controlled crosses between these varieties, and a wild Midwestern population of C. americana sourced from the Wisconsin Department of Natural Resources (DNR) and planted in Barneveld, WI, was sampled following budbreak in May of 2020. Genomic DNA was extracted and libraries for GBS were prepared using a double digestion with the restriction enzymes Nsil and Bfal following the methodology described by Elshire et al. (2011). This specific double digest was selected based on analysis of k-mer distributions of sequence libraries generated with ApeKI alone, Nsil and Mspl, Pstl and Bfal, Pstl and Mspl, ApeKl and Bfal, and ApeKl and Mspl. Illumina GBS barcodes and adapters were ligated, and paired-end reads (2 \times 150 bp, 10 million reads/sample) were generated using an Illumina NovaSeg 6000.

Due to the high levels of synteny across the genomes described above, direct use of Illumina sequence data to infer variable degrees of interspecific hybridization, using tools such as sppIDer (Langdon et al., 2018), is limited. Biallelic SNPs were, therefore, identified using the TASSEL GBSv2 pipeline (Bradbury et al., 2007) and filtered for missing data (<10% across all samples), minor allele frequency (>0.05), linkage disequilibrium ($r^2 < 0.75$) and allele depth (80% of samples with a depth >8) using bcftools (Danecek et al., 2021). Population structure analysis using principal components analysis (Price et al., 2006) has been shown to be a simple and efficient alternative to more complex modelbased approaches such as STRUCTURE (Falush et al., 2007) and ADMIXTURE (Alexander et al., 2009). This filtered VCF file was, therefore, converted to a Euclidean distance matrix using R (R Core Team, 2021), and multi-dimensional scaling (which permits some missing data) was then performed to obtain measures analogous to scores along the first two principal components of the distance matrix.

Comparative analyses of Corylus

Comparisons across the *Corylus* genus were made utilizing the five currently available chromosome-scale genome assemblies, as well as the *C. americana* assemblies reported here for 'Rush' and 'Winkler'. This included two *C. avellana* assemblies (cultivars 'Tombul' (Lucas *et al.*, 2021) and 'Tonda Gentile della Langhe' (Pavese *et al.*, 2021)), two *C. heterophylla* assemblies (an accession from Siping City, Jilin (Liu *et al.*, 2021)) and one *C. mandshurica* assembly (a wild specimen from Xinglong (Li *et al.*, 2021)). All of these assemblies were annotated using similar *ab initio* prediction methods which combined RNAseq data and protein homology data, and thus their transcriptomes were well-suited to comparative analyses with 'Rush' and 'Winkler'.

GENESPACE v0.94 (Lovell *et al.*, 2022) was used to investigate genome-wide syntenic relationships, with *Betula pendula* (silver birch) and *Malus domestica* (apple) included as outgroups (gene models being obtained from https://genomevolution.org/coge/ and https://www.rosaceae.org, respectively). In brief, GENE-SPACE utilizes the program OrthoFinder (Emms and Kelly, 2019) to identify orthogroups from predicted gene models and then parses orthologues to define syntenic blocks across species using BLAST and MCScanX (Wang *et al.*, 2012). Genes in *C. americana* with no orthologous sequences in any other *Corylus* species were also identified using OrthoFinder.

Phylogenetic analysis of sporophytic self-incompatibility (SSI)

Sequences for three S-locus genes (MIK3 homologues) reported by Hou *et al.* (2022) were obtained from the transcriptome reported in Zhao *et al.* (2021). NCBI-BLAST v2.6.0+ (Altschul *et al.*, 1990) was used to identify homologous sequences in all *Corylus* species, as well as *B. pendula*, *M. domestica* and *Brassica oleracea*. These latter three species were included as variably related outgroups which also exhibit self-incompatibly: SSI in the case of *B. oleracea* and *B. pendula* and GSI in the case of *M. domestica*. The *B. oleracea* transcriptome was obtained from http://brassicagenome.net/. Multiple sequence alignments were made using muscle v5.1 (Edgar, 2021). These 84 alignments were then imported into MEGA11 (Tamura *et al.*, 2021) and used to build a phylogenetic tree using maximum likelihood and the Jones-Taylor-Thornton matrix-based model (Jones *et al.*, 1992). A bootstrap consensus tree was inferred from 100 replicates, in which branches present in <50% of these replicates were collapsed. Trees were visualized and clades labelled using the program FigTree v1.4.4.

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Conflict of interest

All authors declare no conflict of interest with the research and findings reported here.

Author contributions

SHB and JCD carried out conceptualization, methodology design and project administration. SHB, DMS and TB performed data curation. SHB, DMS, TB, SS and JCD conducted formal analysis. SHB performed visualization. SHB, JCD and TB wrote the manuscript. SHB and JCD were responsible for funding acquisition.

Data availability statement

Genome assemblies, as well as short- and long-read sequence data (from Arima Hi-C, Oxford Nanopore, PacBio and RNAseg libraries) are available through the NCBI Genome & Sequence Read Archive, respectively, under BioProject ID: PRJNA939214 BioSample accessions: SAMN33458638 ('Rush'). SAMN33458639 ('Winkler'). Genome assemblies, GFF3 models of predicted genes, functionally annotated peptide sequences, anchored pangenome and VCF of GBS-derived SNPs are also available at https://doi.org/10.5281/zenodo.7439335. Finally, genomes and annotations have been added to Phytozome: https://phytozome.jgi.doe.gov/info/Camericanavar_winkler_v1_1 Winkler): https://phytozome.jgi.doe.gov/info/ (for and Camericanavar_rush_v1_1 (for Rush). Bioinformatic pipelines for genome assembly and polymorphism identification are published at: https://github.com/shbrainard/Camericana_pipelines.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Figure S1 Clockwise from top left: the typical involucres, growth habit and nuts of *C. americana* 'Rush', held at the NCGR in Corvallis, Oregon.

Figure S2 Circular plot of genomic features in 'Winkler'. From the outside in: ideogram of the 11 chromosomes of *C. americana*; total gene content, as identified by gene prediction and annotation; transposable elements, as identified by RepeatModeler; heatmap of tandem repeats; line graph of tRNA content.

Figure S3 Pairwise dotplots visualizing pairwise BLAST hits between *C. americana* 'Winkler', and each of the other eight genomes included in the GENESPACE analysis.

File S1 Gene IDs for a subset of the *C. americana* transcriptome consisting of 66 predicted single copy orthologs present in both 'Rush' and 'Winkler', but absent in all other analyzed genomes.