RESEARCH PAPER

Characterizing 3D inflorescence architecture in grapevine using X-ray imaging and advanced morphometrics: implications for understanding cluster density

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Received 28 February 2019; Editorial decision 13 August 2019; Accepted 21 August 2019

Abstract

Inflorescence architecture provides the scaffold on which flowers and fruits develop, and consequently is a primary trait under investigation in many crop systems. Yet the challenge remains to analyse these complex 3D branching structures with appropriate tools. High information content datasets are required to represent the actual structure and facilitate full analysis of both the geometric and the topological features relevant to phenotypic variation in order to clarify evolutionary and developmental inflorescence patterns. We combined advanced imaging (X-ray tomography) and computational approaches (topological and geometric data analysis and structural simulations) to comprehensively characterize grapevine inflorescence architecture (the rachis and all branches without berries) among 10 wild Vitis species. Clustering and correlation analyses revealed unexpected relationships, for example pedicel branch angles were largely independent of other traits. We identified multivariate traits that typified species, which allowed us to classify species with 78.3% accuracy, versus 10% by chance. Twelve traits had strong signals across phylogenetic clades, providing insight into the evolution of inflorescence architecture. We provide an advanced framework to quantify 3D inflorescence and other branched plant structures that can be used to tease apart subtle, heritable features for a better understanding of genetic and environmental effects on plant phenotypes.

Keywords: 3D architecture, inflorescence, morphology, persistent homology, phylogenetic analysis, topological data analysis, Vitis spp., X-ray tomography.

Introduction

Inflorescences are a major feature of the angiosperm lineage whose architectural variation affects fertilization, fruit development, dispersal, and crop yield (Wyatt, 1982; Hake, 2008; de Ribou et al., 2013; Kirchoff & Claßen-Bockhoff, 2013; Périlleux et al., 2014; Chanderbali et al., 2016). These branched reproductive structures with multiple flowers reflect...
the extraordinary diversity across angiosperm species, from an ear of corn to palms with inflorescences measuring 5 m long (Hodel et al., 2015). Yet seemingly simple processes give rise to these vastly different shapes—during development reproductive meristems may either switch to floral identity or proliferate additional inflorescence meristems and branches (Prusinkiewicz et al., 2007). However, such complex topologies have proven difficult to quantify with conventional tools.

Detailed descriptions of inflorescences by trained experts are often unique to specific research communities or groups of taxa, and are not always readily transferable, hindering meaningful comparative analysis (Endress, 2010). Inflorescences are sometimes described typologically: indeterminate or determinate, simple or compound, as a raceme, cyme, panicle or spike, etc. (Wyatt, 1982; Weberling, 1992). Other approaches describe qualitative attributes of inflorescences such as the presence or absence of certain structures, such as secondary branches or bracts (Weberling, 1992; Doebely et al., 1997; Feng et al., 2011; Hertweck & Pires, 2014). A third method for characterizing inflorescences is through quantification of component structures (e.g. branch length, inflorescence length and width, angular traits; Kuijt, 1981; Marguerit et al., 2009; Landrein et al., 2012; Le et al., 2018). Although these classical quantitative approaches facilitate comparative statistical analyses, the three–dimensional (3D) complexity of inflorescences is largely undescribed. Furthermore, descriptions may be confounded by developmental stage at the time of measurement, and distinguishing between vegetative and reproductive measures is largely undescribed. Furthermore, descriptions may be confounded by developmental stage at the time of measurement, and distinguishing between vegetative and reproductive branching structures can be difficult (Wyatt, 1982; Weberling, 1992; Guédon et al., 2001). Thus, new technological and analytical approaches that can represent comprehensive, multidimensional information about inflorescence diversity are needed to normalize and enrich analysis of these structures.

One promising approach for capturing 3D shapes of inflorescences and other plant structures is X-ray tomography (XRT). XRT generates high quality reconstructions of the internal and external shapes of plants, preserving nearly complete geometric and topological information in 3D. These 3D digital models then can be used to extract quantitative data (features) from plant structures. X-rays have been used to quantify wheat and rice seed structure of barley seedlings (Pfeifer, 2018), and dynamic starch accumulation in living grapevine stems (Earles et al., 2018), among others. Most critically, whereas manual measurements can be laborious and destructive, non-destructive sampling for XRT analysis facilitates comprehensive quantification of complex morphological traits.

Quantifying complex shapes with XRT requires appropriate analytical approaches. Topological modeling, a mathematical field concerned with the connectedness of branching structures, can quantify inflorescence architecture by parsing geometric 3D structures into distinct, yet connected, components (Godin & Caraglio, 1998). Topological modeling has yielded important insights into inflorescence development, functional analysis, and crop improvement in a variety of plant species (e.g. Arabidopsis thaliana, Capsicum annuum, Malus pumila, and Triticum; Godin et al., 1999; Letort et al., 2006; Kang et al., 2009). While powerful, these reductionist approaches rely on an a priori understanding of the mechanisms that contribute to complexity (e.g. branching patterns), and lose power when shapes vary drastically from one another (e.g. comparing a corn tassel to a grape cluster). Approaches that capture emergent properties of complex structures without presupposing the importance of individual structural components are complementary to traditional topological models (Bucksch et al., 2017).

An emerging mathematical approach to interpret topological models is persistent homology (PH). PH extracts morphological features from two- or three-dimensional representations and can be used to compare very different shapes. PH has been applied to explain a wide range of features including atomic structures, urban and forested areas, cancers, cell shapes, and jaw shape, among others (Edelsbrunner & Morozov, 2013). In plants, PH has been used to estimate shapes that are otherwise difficult to measure including leaves, leaflet serration, spikelet shape, stomatal patterning, and root architecture (Haus et al., 2018; Li et al., 2018a,b; Migicovsky et al. 2017; McAllister et al., 2019). Previous work showed that PH could capture more quantitative variation than traditional plant morphological measures (described above) resulting in the identification of otherwise latent quantitative trait loci (Li et al., 2018b). PH is especially well-suited for quantifying branching topology as it can quantitatively summarize complex variation with a single measure (Li et al., 2017; Delory et al., 2018). Rachis, pedicel, and branches include inherently topological features that can be especially well-analysed with PH-based methods.

Grape clusters (or bunches) are branched structures supporting berries produced by grapevines (Vitis spp.) and are an ideal system in which to apply XRT and PH. Grape inflorescences are historically, culturally, and economically important and vary extensively in nature and in cultivation (Iland et al., 2011). Cluster architecture determines bunch density, which is described as ‘arrangement of berries in a cluster and the distribution of free space’ (Richter et al., 2019). The density of berries in a cluster is an important breeding feature because it determines yield, wine character, and disease resistance (amount of air flow between berries is a primary determinant of pests and pathogens on the fruit). Cluster density is a characteristic identified by the Organization Internationale de la Vigne et du Vin, and varies from ‘berries clearly separated’ (loose clusters) to ‘berries deformed by compression’ (very dense clusters; OIV, 2001). As primary determinants of yield, end-product characteristics and disease resistance cluster architecture have been studied extensively in grapevine (reviewed in Tello & Ibáñez, 2018). These studies have shown that wine grape cultivars (Vitis vinifera) display distinct bunch densities (Shavrukov et al., 2004). However, less is known about cluster architecture in wild Vitis species, an important source of natural variation used by breeders in the development of hybrid grapevine varieties.

Historically, researchers have focused on a suite of cluster traits such as cluster size, shape, weight, and density/
compactness to characterize bunch density quantified in grapevines (Rovasenda, 1881; Pulliat, 1888; Bioletti, 1938; Galet, 1979; Bettiga, 2003). Measurements are made primarily using traditional tools including rulers, digital calipers, volume displacement, and/or through human judging panels. More recently, automated image-based approaches have been implemented to capture aspects of cluster architecture in the lab and field (Ivorra et al., 2015; Aquino et al., 2017, 2018; Rist et al., 2018). However, these image-based methods cannot penetrate the internal inflorescence structure. Therefore resulting models are based only on the visible surface and the underlying topology cannot be fully captured, limiting an understanding of how inflorescence architecture and berry features co-vary. XRT and PH applications offer an important opportunity to understand grapevine bunch density through detailed analyses of inflorescence architecture. This work will deepen our understanding of natural variation of inflorescence structure, identify priority targets for breeding, and permit connecting 3D structure to underlying processes and genetics of inflorescence development.

We used X-ray tomography, geometric measurements, persistent homology, and structural simulation to characterize wild grapevine inflorescence architecture. We targeted the branching architecture of the mature inflorescence: the rachis and all branches that remain following the removal of ripe berries (Fig. 1). Specifically, we aimed to: (i) characterize variation in component traits of inflorescence architecture within and among Vitis species; (ii) assess phylogenetic signals underlying inflorescence architecture traits; and (iii) interpret inflorescence trait variation in the context of breeding objectives. This work represents an important advance for the characterization of 3D plant architecture using a powerful combined imaging and computational approach.

Materials and methods

Plant material

In this study, we sampled grapevine bunches from 136 unique Vitis species representing 10 wild Vitis species living in the USDA germplasm repository system (Geneva, NY, USA; Table 1; Supplementary Fig. S1 at JXB online). Grapevines have a paniculate inflorescence that consists of a rachis with several primary and secondary branches, tapering towards the terminus of the organ (Iland et al., 2011). Wild grapevines are dioecious; consequently, unbalanced sample sizes for different species reflect numbers of female genotypes available in the germplasm collection. Each unique genotype is represented in the germplasm collection by two clonally replicated vines. For most of the 136 genotypes, we collected a total of three clusters from the two clonal replicates combined, representing average cluster morphology. We avoided clusters that were visibly damaged or indirectly altered (e.g. tendril or trellis interference). For each vine, clusters were removed from separate canes at the point of peduncle attachment (Fig. 1A). In total, 392 clusters were collected in September 2016 when berries were soft, equivalent to EL.38 developmental stage (Coombe, 1995; Fig. 1A). Berries were manually removed from clusters in the field, and the remaining inflorescence stalks (including rachis, branches, and pedicels; hereafter referred to as inflorescence or inflorescence architecture) were used to assess inflorescence architecture (Fig. 1B).

X-ray tomography and data preprocessing

Grapevine inflorescences were scanned at the Donald Danforth Plant Science Center (St Louis, MO, USA) using a North Star Imaging X5000 X-ray tomography instrument (NSI; Rogers, MN, USA) equipped with a 16-bit Varian flat panel detector (1536×1920 pixels with 127 µm pixel pitch) and 225 kV microfocus reflection target X-ray source. Each inflorescence was held between two pieces of construction-grade expanded polystyrene, clamped in a portable clamping device, and positioned on the X-ray turntable in one of two configurations (Fig. 1C): 725 mm from the source, generating ×1.26 magnification and 107 µm voxel resolution, or 766 mm from the source, generating ×1.19 magnification and 107 µm voxel resolution. Each scan used X-ray power set to 60 kV and 1200 µA at 10 frames per second, collecting 1200 16-bit TIFF projections over 360° of rotation during a 2 min continuous standard scan. Projections for each scan (Fig. 1D) were combined into a single 3D volume using NSI eX-CT software, converted to a density-based surface rendering.
Table 1. Number of samples/individuals of each species and berry information used in the study

| Species              | Number Samples | Individuals   | Individuals used in phylogenetic analysis |
|----------------------|----------------|--------------|-------------------------------------------|
| V. acerifolia        | 32             | 11           | 9                                         |
| V. aestivalis        | 5              | 2            | 1                                         |
| V. amurensis         | 13             | 5            | 2                                         |
| V. cinerea           | 45             | 15           | 13                                        |
| V. coignetiae        | 6              | 2            | 1                                         |
| V. labrusca          | 62             | 22           | 12                                        |
| V. palmata           | 3              | 1            | 1                                         |
| V. riparia           | 158            | 53           | 48                                        |
| V. rupestris         | 41             | 16           | 10                                        |
| V. vulpina           | 27             | 9            | 2                                         |
| Total                | 392            | 136          | 99                                        |

| Berry information*  | Low diameter (mm) | High diameter (mm) | Berries per bunch |
|---------------------|-------------------|--------------------|-------------------|
| V. acerifolia       | 8                 | 12                 | >25               |
| V. aestivalis       | 8                 | 20                 | >25               |
| V. amurensis        | 8                 | 15                 | NA                |
| V. cinerea          | 4                 | 8                  | >25               |
| V. coignetiae       | NA                | 8                  | NA                |
| V. labrusca         | 12                | 23                 | <25               |
| V. palmata          | 8                 | 10                 | >25               |
| V. riparia          | 8                 | 12                 | >25               |
| V. rupestris        | 8                 | 12                 | <25               |
| V. vulpina          | 8                 | 12                 | >25               |

* Berry information is from Galet (1988) and Moore and Wen (2016).

Polygon file (PLY), and exported for analysis (Fig. 1E). The full PLY dataset for this work is 7.85 GB, and can be downloaded from: https://www.danforthcenter.org/scientists-research/principal-investigators/chris-topp/resources.

We exported the surface mesh data (.ply files) into Meshlab (v1.3.3; Cignoni et al., 2008) and performed the following processing steps to remove topological noise: (i) deleted the vertices where branches touch using ‘Select Vertices’ and ‘Delete Selected Vertices’ filters; (ii) removed duplicates and isolated vertices and faces using the filters ‘Remove Duplicated Vertex’, ‘Remove Duplicate Faces’, ‘Remove Isolated pieces (wrt Diameter)’, and ‘Remove Unreferenced Vertex’.

**Geometric inflorescence architecture traits**

We extracted 15 geometric traits from scanned inflorescences (Fig. 2; Supplementary Fig. S2). Detailed trait descriptions and calculations are explained in Supplementary Table S1. Trait illustrations, including examples of low and high values for each trait, are available in Fig. 2 and Supplementary Fig. S2. Traits were organized in one of three trait groups: global-size features, local-branching features, and size-invariant features (Table 2). Pedicel Diameter and Pedicel Branch Angle were measured using the software DynamicRoots (Symonova et al., 2015) on a subset of detected pedicels from the raw 3D volume data. All other traits were derived from Matlab algorithms. Branch length traits (i.e. Total Branch Length, Rachis Length, Pedicel Length, and Avg Branch Length) were derived from the persistence barcode (see next subsection).

**Quantifying branching topology using persistent homology, a topological data analysis method**

Persistent homology measures shapes based on a tailored mathematical function, such as geodesic distance, which we used here to capture both curved length and topology of the branches (Fig. 3; Supplementary Video S1). The geodesic distance of a point is the length of the shortest curve connecting the point and the base (e.g. purple curves, Fig. 3A), where the tailored base can be set as the first node or ground level (the brown line in Fig. 3A). For each branch, the tip always has the largest geodesic distance from the base (Fig. 3B). A level represents the collection of points whose geodesic distances are the same (e.g. geodesic distance = 90, pink curve in Fig. 3A). A superlevel set, for example, at 90, is all the points whose geodesic distances are greater than 90 (black branch tips, Fig. 3A). Changing the level value from largest to smallest (x-axis, Fig. 3C), the sequence of nesting superlevel sets can be formed, which is named superlevel set filtration (top panel, Fig. 3C). During the change of the level value, bars record the connected components for each of the superlevel sets. When a new component arises, a new bar starts (e.g. at level 112, purple branch, Fig. 3C). When two components merge (e.g. at level 65, orange branch merges into purple branch, Fig. 3C), the shorter bar stops (e.g. the orange bar stops at level 65, Fig. 3C). This bar graph, called the persistence barcode, summarizes topological information such as branching hierarchy, branch arrangement, and branch lengths. In our study, we set the base as the junction between peduncle and rachis (the lowermost node, indicated by a brown line in Fig. 1E and Fig. 3D,F) and use this base to compute the persistence barcode for the inflorescence architecture (Fig. 3E, G).

The persistence barcode can be used to compare topological similarity between any two inflorescences. To compute pairwise distance among persistence barcodes for the entire inflorescence population, we used the bottleneck distance (Cohen-Steiner et al., 2007). Bottleneck distance is a robust metric that calculates the minimal cost to move bars from one persistence barcode to resemble another (Li et al., 2017). We performed multidimensional scaling (MDS) on the pairwise bottleneck distance matrix and projected the data into lower dimensional Euclidean space by preserving the pairwise distance as well as possible. The Matlab (R2017a) MDS function cmdscale() projects the data so that MD1 acts as principal component (PC) 1 representing the most variation. The first three PCs (MDs) explained about 80% of the total variation and were included as traits: PersistentHomology_PC1 (PH_PC1, explained about 54% of the variation), PersistentHomology_PC2 (PH_PC2, explained about 20% of the variation), and PersistentHomology_PC3 (PH_PC3, explained about 6% of the variation). Those traits not only measure the topological structure, but also relate to geometric variation (e.g. global size) as the data were not normalized (Fig. 2; Supplementary Table S1).

Next, we normalized the persistence barcode by the Total Branch Length (summation of the bar lengths) so that the Total Branch Length was 1. By a similar procedure, we derived the first three PCs named PersistentHomologyNormalizedByTotalBranchLength_PC1 (PHn_PC1, explained about 45% of the variation), PersistentHomologyNormalizedByTotalBranchLength_PC2 (PHn_PC2, explained about 21% of the variation), and PersistentHomologyNormalizedByTotalBranchLength_PC3 (PHn_PC3, explained about 7% of the variation) for the normalized inflorescence topological structure (Fig. 2; Supplementary Table S1).

**Berry potential, an approach to indirectly explore the space limited by inflorescence architecture**

An ongoing question in grapevine cluster architecture is the relationship between inflorescence architecture and berry number and size. Inflorescence architecture is one of several factors determining the number of berries that can form, due to the number of pedicels and the available space for berry development. In this study, berries were removed because of concerns about berry integrity during transport from New York to Missouri, and the time between harvest and scanning. Instead of manually counting the number of berries per bunch, we used the persistence barcode to estimate berry potential, which is a robust metric that calculates the minimal cost to move bars from one persistence barcode to resemble another (Li et al., 2017). We performed multidimensional scaling (MDS) on the pairwise bottleneck distance matrix and projected the data into lower dimensional Euclidean space by preserving the pairwise distance as well as possible. The Matlab (R2017a) MDS function cmdscale() projects the data so that MD1 acts as principal component (PC) 1 representing the most variation. The first three PCs (MDs) explained about 80% of the total variation and were included as traits: PersistentHomology_PC1 (PH_PC1, explained about 54% of the variation), PersistentHomology_PC2 (PH_PC2, explained about 20% of the variation), and PersistentHomology_PC3 (PH_PC3, explained about 6% of the variation). Those traits not only measure the topological structure, but also relate to geometric variation (e.g. global size) as the data were not normalized (Fig. 2; Supplementary Table S1).

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of looking directly at berries on the cluster, we used inflorescence architecture as a starting point to simulate potential space available for berry growth by evaluating expanding spheres attached to pedicels. The extent of sphere expansion allowed by each pedicel is referred to as ‘berry potential’ (Fig. 4; Supplementary Video S2).

We first determined the growth direction for each berry potential based on the pedicel orientation. When spheres expand, the center moves along the pedicel direction (Fig. 4A). This step can be achieved by performing principal component analysis (PCA) on the near-berry segment of the pedicel. The first principal axis is the pedicel direction. We adjusted the arrow of the direction to make sure berry potential increases outward along the pedicel orientation. Then the berry potential increases until one of three situations is encountered (Fig. 4B): (i) if two berry potentials touch each other, both berry potentials will stop increasing; (ii) if a berry potential touches any part of the inflorescence, it will stop increasing; (iii) if the diameter of the berry potential reaches the maximum size known for that species (Table 1), it will stop increasing. For each species, the maximum size is defined as the maximum berry diameter, a number estimated from known ranges of berry sizes for each species, based on values obtained from Galet (1988) and Moore and Wen (2016).
Berry potential does not reflect true berry growth; rather, berry potential is a derived attribute of inflorescence architecture, an indirect estimate of the space potentially available for berry growth. It also does not account for the possibility of branches bending or otherwise becoming reoriented due to pressure from growing berries. Berry potential is based on the number of neighbor pedicels, neighbor pedicel lengths, and neighbor pedicel mutual angles. Larger values for berry potential are associated with fewer neighbor pedicels and/or longer pedicel lengths and/or larger mutual angles. From the berry potential simulation, we calculated three features, TotalBerryPotentialVolume, AvgBerryPotentialDiameter, and BerryPotentialTouchingDensity, which is the berry potential touching number (i.e. touching either another berry potential or any part of the inflorescence) divided by the number of berry potential (Fig. 2; Supplementary Table S1).

To compare the berry potential with the true berry features, we XRT scanned 10 clusters with berries attached representing five commercially available table grape (*Vitis vinifera*) varieties. We computationally identified the berries and measured the total berry volume, average berry diameter, and touching density. We then computationally removed the berries and calculated the berry potential features based on the stalk. The linear regression ($R^2=0.93, 0.99, $ and 0.76, respectively) and normalized root mean square error (RMSEn=11, 3, and 14, respectively) indicate that berry potential provides a realistic model of actual grapevine cluster traits (see Supplementary Fig. S3). We note that the current simulation will better estimate relatively loose clusters with round berries than highly compact clusters in which berries touch, are deformed by pressing, and are not spherical. Our intent is to further parameterize the simulation by scanning even more diverse germplasm to incorporate a wider range of cluster architectures and berry traits in future work.

**Phylogenetic analysis**

Phylogenetic analyses were conducted to understand evolutionary trends in inflorescence architecture in *Vitis*. Single nucleotide polymorphism (SNP) markers were generated as part of a separate study of the USDA Grapevine Germplasm Reserve in Geneva, NY, USA (Klein et al., 2018). The original dataset consisted of 304 individuals representing 19 species that were sequenced using genotyping-by-sequencing (GBS; Elshire et al., 2011). Briefly, Klein et al. (2018) filtered data to retain biallelic sites with a minimum allele frequency of 0.01, a minimum mean depth of coverage of 10×, and only sites with <20% missing data and individuals with <20% missing data. We extracted SNP data for 99 of the individuals from this study that were also genotyped in Klein et al. (2018; Table 1) and performed phylogenetic analysis using SVDquartets (Chifman & Kubatko, 2014), a maximum likelihood approach designed to address ascertainment bias associated with reduced representation sequencing techniques like GBS. We analysed all possible quartets and carried out 100 bootstrap support runs (see Supplementary Fig. S1) using PAUP* version 4.0a (Swofford, 2003). The three main clades recovered in the tree were consistent with previous phylogenetic work in *Vitis*: (i) an Asian clade (*V. amurensis* and *V. oigetica*), (ii) North American clade I (*V. riparia, V. acerifolia, and V. rupestris*), and (iii) North American clade II (*V. vulpina, V. cinerea, V. aestivalis, V. labrusca, and V. palmata*) (Tröndle et al., 2010; Zecca et al., 2012; Miller et al., 2013; Zhang et al., 2015; Klein et al., 2018).

To visualize trait distributions on a phylogenetic tree using branch lengths, we used Mega X (Kumar et al., 2018) to generate a neighbor joining tree with 2000 bootstrap replicates. All measurements were averaged across the three replicates per genotype to produce an average value for each trait for each genotype. We computed Pagel’s lambda to estimate phylogenetic signal for each morphological trait and mapped each trait onto the phylogeny (see Supplementary Fig. S4A–X) using the R package phytools (v. 0.6–44; Revell, 2012). We calculated variation of each morphological trait for each clade based on the mean value for each species (Supplementary Fig. S5).

**Statistical analysis**

PCA, MDS, and hierarchical cluster analysis generating a hierarchical tree were performed in Matlab using functions pca(), cmdscale(), and clustergram(). The R function cor.mtest() and package corrplot (Wei & Simko, 2017) were used for significance tests and correlation matrix visualization. The function lda() in R package MASS (Venables & Ripley, 2002) was used for the linear discriminant analysis (LDA) with a jack-knifed ‘leave one out’ cross validation method.

**Code availability**

All Matlab functions used to calculate persistence barcodes, bottleneck distances, simulation for berry potential, other geometric features used in this study, and the script for extracting phylogenetic information can be found at the following GitHub repository: https://github.com/Topp-Roots-Lab/Grapevine-inflorescence-architecture.

**Results**

**Inflorescence morphological variation and trait correlation within Vitis species**

We investigated 24 morphological traits (15 geometric traits, six PH traits, and three berry potential traits) of inflorescence architecture in 10 wild *Vitis* species (136 genotypes, 392 samples) and detected wide variation in morphological features within and between species (Fig. 2; Supplementary Fig. S2; Supplementary Table S2). In particular, of all the species examined, *V. aestivalis* had the largest variance for TotalBerryPotentialVolume. *Vitis labrusca* had the largest variance for 10 traits (i.e. pedicel features, Sphericity, AvgBranchDiameter, AvgBerryPotentialDiameter, and normalized topological traits). *Vitis cinerea* had the largest variance for six traits (i.e. most global-size features, PH_PC2, and PH_PC3). In comparison, *V. palmata* had the smallest variance for eight traits (i.e. pedicel features, Sphericity, AvgBranchDiameter, TotalBerryPotentialVolume, PH_PC3, and PHn_PC3), as did *V. amurensis* (global-size features, RachisLength, PH_PC1, and PH_PC2).

All traits were hierarchically clustered based on the mean trait values for each species, classifying traits into two main categories: mostly size-invariant+local-branching features (PHn_PC3 to PedicelLength) versus global-size features (AvgBranchLength to BerryPotentialTouchingDensity) (Fig. 5A). Hierarchical clustering (Fig. 5A) and pairwise correlation for morphological traits (Fig. 5B) showed that global-size features (ConvexHullVolume, SurfaceArea, Volume, NumberOfPedicel, and TotalBranchLength), PH_PC1, and RachisLength were all highly positively correlated. We refer
to these seven traits as size-associated features. Size-associated features were negatively correlated with PedicelLength/RachisLength, Solidity, Sphericity, and PHn_PC1. Some traits were relatively independent such as 2nd/LongestBranchLength, PedicelLength, PedicelBranchAngle, PH_PC2, PHn_PC2, and PHn_PC3 (Fig. 5B). PH_PC3 had some negative relation with size-invariant features. PHn_PC1 positively correlated with Sphericity, Solidity, and AvegeBerryPotentialDiameter (Fig. 5B).

Pairwise correlations of morphological features (allometric relationships) for each of the species varied widely (Fig. 5C; for all traits see Supplementary Fig. S6A–X). For example, more pedicels typically result in smaller berry potential diameters, except for V. aestivalis. Longer branches tended to be thinner, except for V. coignetiae, and correlate with larger inflorescences, except in V. acerifolia.

Hierarchical clustering of 10 Vitis species based on the 24 morphological traits resolved four groups: (i) V. cinerea, (ii) V. aestivalis, (iii) V. coignetiae–V. vulpina–V. palmata–V. acerifolia–V. riparia–V. rupestris, and (iv) V. amurensis–V. labrusca (Fig. 5A). Among the 10 Vitis species examined in this study, the largest variance in mean trait values were seen in V. cinerea (Fig. 5A). Vitis cinerea samples were generally larger than those from...
the other species, as reflected in size-associated traits. Topology traits such as PHn_PC3 and size-invariant traits like Sphericity and Solidity were lower in the mean trait value for V. cinerea than for other species. Similarly, mean trait values were larger for size-associated traits in V. aestivalis (Fig. 5A). Compared with other species, topology and berry potential traits were larger in V. aestivalis. Mean trait values of the third group (V. coignetiae–V. vulpina–V. palmata–V. acerifolia–V. riparia–V. rupestris, Fig. 5A) tended to be nearer to middle values compared with the other species. Within this group, V. acerifolia–V. riparia–V. rupestris typically were larger in the mean trait value for berry potential touching (i.e. denser berry potentials). These three species and V. palmata tend to have large, first primary branches (i.e. wings; Fig. 1E). Vitis coignetiae has thicker branches and V. vulpina has longer pedicels compared with other species in this group. The final group, V. amurensis and V. labrusca, have relatively smaller inflorescences with thicker branches compared with the other species sampled here. These general features are reflected in larger mean values for several size-invariant and local-branching features and smaller mean values for many branch length-dependent and size-associated features, respectively (Fig. 5A).

**Multivariate, discriminant analysis of Vitis species based on inflorescence architecture**

In order to understand how overall inflorescence architecture varies among Vitis species, we performed PCA using all 24 morphological features and all samples. PC1 explained 37.12% of the total variation in the measured architecture (Fig. 6A). The traits with the largest values for PC1 loadings, indicating that they contributed most to variation, were size-associated features, Solidity and Sphericity. PC2 explained 15.4% of the total variation in the measured inflorescence architecture, with variation primarily explained by local-branching features such as PedicalDiameter, PedicelLength, PedicelLength/RachisLength, AvgBranchLength, BranchDiameter, three berry potential traits, and PHn_PC1 (Fig. 6A). Although inflorescences from each species occupy different regions of morphospace, these regions overlap considerably.

LDA performed on the first 18 PCs explaining 99.5% of the variation, distinguished between species with a classification accuracy rate of 78.32%. A confusion matrix (Fig. 6B) shows the proportion of samples correctly predicted for each species. LD1 primarily separates V. cinerea, V. labrusca, and V. amurensis from the other species while LD2 primarily separates V. vulpina and V. coignetiae. The traits that are most important for distinguishing these species, as indicated by LD loadings, are TotalBerryPotentialVolume and PHn_PC1 for LD1, and AvgBranchLength and AvgBerryPotentialDiameter for LD2 (Fig. 6B). The most important predictors for correctly separating any two species are shown as the grey scaled boxes in Supplementary Fig. S7 and Table S3. For example, BranchDiameter and PedicelDiameter are key when contrasting V. coignetiae and V. vulpina, suggesting that different branch thickness easily distinguishes these two species. This method correctly determined species classifications with 100% accuracy when contrasting V. aestivalis and V. cinerea, V. aestivalis and V. palmata, V. aestivalis and V. vulpina, V. amurensis and V. cinerea, V. amurensis and V. palmata, V. cinerea and V. coignetiae. Other combinations of species are harder to distinguish on the basis of inflorescence characters. For example, the classification accuracy rate was only 80% when distinguishing between V. amurensis and V. labrusca and 82% for V. aestivalis and V. coignetiae.

**Phylogenetic signal of inflorescence architecture within clades**

The phylogeny dataset (n=99) is generally well-supported at the species level and correlates well with current taxonomy. Using average trait values per individual, Pagel's lambda showed 12 morphological traits (seven size-associated features along with PedicalDiameter, TotalBerryPotentialVolume, Sphericity, PH_PC2, PHn_PC1) have a strong phylogenetic signal (lambda>0.8; Fig. 7; Supplementary Table S4). While most species sampled tended to have small values for the seven size-associated features, V. aestivalis, V. cinerea, and V. vulpina tended to have values that were either close to median or larger. On average, V. labrusca had larger values for Sphericity and PHn_PC1 compared with other species sampled, while V. cinerea generally had some of the smallest values for these traits. Only two morphological traits (2nd/LongBranchLength, lambda=0.06 and BerryPotentialTouchingDensity, lambda=0.25) lacked phylogenetic signal (Fig.7; Supplementary Table S4).
We observed differences in *Vitis* inflorescence architecture among clades and between species. For North American (NA) clade I (*V. acerifolia*, *V. riparia*, *V. rupestris*), variation in the 24 morphological traits measured had similarly small values among species, particularly for several size-associated traits, although there was relatively large variation for PH_PC3 and BerryPotentialTouchingDensity (Fig. 7). Within NA clade I, we observed differences among clade members for traits such as Sphericity and PHn_PC1 (larger in *V. rupestris* compared with other clade members) and PedicelDiameter and BranchDiameter (slightly larger in *V. acerifolia* compared with other clade members; Fig. 7). NA clade II appeared to be more variable among clade members. *Vitis cinerea* had larger values for size-associated traits compared with clade members *V. labrusca*, *V. palmata*, and *V. vulpina*. Meanwhile, *V. labrusca* typically had larger values for local features (e.g. Sphericity, PedicelDiameter, AvgBerryPotentialDiameter, PedicelBranchAngle) compared with the other clade members (Fig. 7).

We calculated the mean value for each species of each morphological trait to study variation within the three clades and detect subtle signatures (Fig. 7). We computed the variance for the multivariate trait (combining all the 24 traits), and each
of these 24 traits for each clade (see Supplementary Fig. S5; Supplementary Table S5). Overall, based on the samples used in this analysis, variance of the multivariate trait for NA clade I (variation=0.14) was much smaller than that for NA clade II (variation=0.64), while the variation for the Asian clade was 0.39. Some traits had almost no variance in the Asian clade such as PedicelDiameter, PHn_PC2, PH_PC3, and 2nd/LongestBranchLength. However, North American species (8/~19 taxa) in this study were better represented than Asian species (2/~37 taxa), so we are cautious not to over-interpret this finding. Traits with the greatest variance in the Asian clade included PedicelLength/RachisLength, RachisLength, and PH_PC1, while NA clade I had the greatest variance in PHn_PC2. All the other traits had greatest variance in NA clade II (Supplementary Fig. S5, Supplementary Table S5). Traits with the smallest variance in the Asian clade included PHn_PC3, PHn_PC1, PedicelDiameter, BranchDiameter, NumberOfPedicel, 2nd/LongestBranchLength, PH_PC3, and BerryPotentialTouchingDensity. The other traits had small variance in NA clade I (Supplementary Fig. S5, Supplementary Table S5). Our results highlight clade-specific variation in inflorescence architecture for previously undescribed traits.
Fig. 7. Phylogenetic analysis. A neighbor joining phylogenetic tree for a subset of the *Vitis* dataset (n=99). Node values denote bootstrap support for values ≥50. Ten *Vitis* species are highlighted in different colored backgrounds. Three clades (Asian clade, NA clade I, and NA clade II) are labeled and marked by vertical bars. The barplot showing values of Pagel's lambda, an estimate of phylogenetic signal, overlaps with the trait name on the right top panel. Below each trait, a rainbow colormap shows the values for individuals (small values in red to large values in blue). Rectangles surround the trait value map for species with more than five individuals. One trait (PHn_PC1) was randomly selected to be projected onto the phylogenetic tree branches, and indicates trait variation (red, lower values; blue, higher values) within individuals and among clades.
Discussion

X-ray imaging and advanced morphometrics is a powerful combination for characterizing complex phenotypes

Inflorescence architecture has a strong influence on flower and fruit production and is therefore a trait of great scientific interest. Studies extend into interspecific variation, pollen dispersal, genetic architecture, evolution, regulation, and development of inflorescence structures (e.g. Bradley et al., 1996; Friedman & Harder, 2004; Kellogg, 2007; Morris et al., 2013; Han et al., 2014; Hodge & Kellogg, 2015; Whipple, 2017; Stitzer & Ross-Ibarra, 2018; Ta et al., 2018; Richter et al., 2019). Yet the challenge remains to analyse these complex 3D branching structures with appropriate tools. High resolution datasets are required to represent the actual structure and comprehensive analysis of both the geometric and topological features relevant to phenotypic variation and to clarify evolutionary and developmental inflorescence patterns.

Our results demonstrate the power and potential of X-ray imaging and advanced morphometric analysis for investigating complex 3D phenotypic features. We analysed the phenotypic variation in inflorescence architecture of 10 wild Vitis species using computer vision and an emerging biological shape analysis method, persistent homology, which allowed comprehensive comparisons of shape. Although samples analysed here represent only a subset of the known variation in Vitis, which includes an estimated 60 species, our analyses demonstrate significant variation within and among Vitis species and among clades. Correlation analysis (Fig. 5B) revealed some unexpected relationships, for example pedicel branch angles were largely independent of other traits. It also shows that PH features are complementary, as they are relatively independent from most geometric features. We were able to assign widely differing architectures to biological species with high accuracy (Fig. 6) from the 24 different morphometric traits surveyed in this study. PH provides an important contribution to this discriminatory power, as does berry potential (Fig. 6B). We observed that traits such as the rachis length, the sum of all branches, the space encompassing the inflorescence architecture (ConvexHullVolume), and PH features can be indicative of species and clade (Fig. 7). Our results suggest meaningful, comprehensive information about the inflorescence structure was captured with a single measure (i.e. the persistence barcode) and that PH is a valuable method for quantifying and summarizing topological information.

Persistent homology analysis has led to a deeper understanding of trait genetic variation and architecture in plants (Li et al. 2018a,b). In grapevine, quantitative trait locus (QTL) analysis indicates a genetic basis to inflorescence architecture and berry compactness (Correa et al., 2014; Richter et al., 2019). Deploying PH-based topological modeling to grapevine mapping populations could lead to the rapid identification of additional inflorescence trait QTLs for breeding. For example, we observed total branch length (a proxy for bigger or smaller clusters) correlates with number of pedicels (a proxy for berry number; Fig. 5), an informative relationship to assess potential yield. However, selecting for total branch length might lead to a negative correlation with the average berry potential diameter (i.e. smaller berries). Although this correlation may be desirable for wine grapes, it is not for table grapes.

Linking complex phenotypes with evolutionary patterns to facilitate precision breeding

Grapevine cluster architecture is a composite feature that reflects multiple subtraits including stalk traits (inflorescence architecture) and berry features (Richter et al., 2019). OIV 204 uses ‘bunch: density’ to describe variation in clusters, ranging from 1, berries clearly separated with many visible pedicels, to 9, berries deformed by compression (OIV, 2001; Rombough, 2002). Other authors have deconstructed traits contributing to cluster architecture primarily through individual measurements collected manually (e.g. Shavrukov et al., 2004; Tello et al., 2015; Zdunić et al., 2015; Tello & Ibáñez, 2018) and more recently, with image-based technologies (Cubero et al., 2014; Roscher et al., 2014; Ivorra et al., 2015; Aquino et al., 2017, 2018; Rist et al., 2018). Here, we were able to describe traits of interest that contribute greatly to the morphological features captured by the OIV scale (e.g. NumberOfPedicel, PedicelLength, PedicelBranchAngle, RachisLength, overall shape using PH; Fig. 2; Supplementary Fig. S2). This method could facilitate precision breeding for both whole inflorescence structure topology and specific desirable geometric traits.

While several studies have quantified cluster structure in cultivated grapevines, similar studies of wild Vitis inflorescence architecture are lacking. Munson (1909) and Galet (1979) describe North American Vitis cluster structure qualitatively, commenting on compactness, size, shape, and the presence of large first primary branches (wings/shoulders). Taxonomic descriptions typically do not examine inflorescence architecture beyond categorical type, position on the vine, and the average number of berries per cluster (Comeaux et al., 1987; Moore, 1991; Moore & Wen, 2016). Descriptions of the position of the inflorescence are useful for identification and are included in dichotomous keys; however, to our knowledge, other inflorescence architecture traits have not been rigorously quantified among wild Vitis species. Although qualitative descriptions are valuable and accessible, powerful phenotyping tools are required to associate complex phenotypes with evolutionary and developmental patterns.

Using 3D imaging and PH with a topological modeling approach, we identified attributes of inflorescence architecture that vary within and among Vitis species that, to our knowledge, have not been previously described. Differences in inflorescence architecture among clades mirror other phenotypic differences among members of North American Vitis. For example, members of NA clade I (V. acerifolia, V. riparia, and V. rupestris) have small values for size-associated features (e.g. RachisLength, ConvexHullVolume, NumberOfPedicel, TotalBranchLength, SurfaceArea, Volume) and relatively large values for PH_PC3 and BerryPotentialTouchingDensity (Fig. 7). These species share suites of other morphological characters (nodal diaphragm, branch and leaf surface traits, and large
stipules; Moore 1991, Moore and Wen 2016, Klein et al., 2018). It is possible that among closely related species conserved pathways generate vegetative and reproductive similarities.

Sample size is low for the Asian clade and most of NA clade II, limiting our ability to assess variation in these species; however, members of NA clade II do not have suites of shared inflorescence traits (V. aestivalis, V. cinerea, V. labrusca, V. vulpina; Klein et al., 2018). Rather, V. labrusca has very small values for size-associated traits and larger values for local features compared with the other clade members, whereas V. cinerea has larger values for size-associated features and smaller values for local features (Fig. 7). This is consistent with the observation that aside from core phenotypic synapomorphies in the genus (tendril, bark, lenticel, and nodal diaphragm characters), members of NA clade IIb (V. aestivalis, V. cinerea, V. labrusca, and V. vulpina) do not share morphological traits unique to the clade (Klein et al., 2018). These species mostly co-occur across their distributions (Callen et al., 2016) and additional sampling of Vitis taxa is necessary to further explore these complex evolutionary patterns. We observed V. amurensis grouping with V. labrusca and V. coignetiae grouping with North American species in hierarchical cluster analysis (Fig. 5A). The former two species have relatively smaller inflorescence architectures with thicker branches compared with the other species sampled here. Taxonomic relationships among North American and Asian Vitis species have been historically challenging, with clades comprising species with disjunct distributions (Mullins et al., 1992). Since current taxonomy resolves separate Asian and North American clades (Klein et al., 2018), morphological similarity between these species likely reflects convergent evolution.

Future directions

Three-dimensional imaging through XRT and advanced mathematical approaches like persistent homology provide new ways to visualize and interpret complex biological structures including inflorescences, and to understand the genetic and environmental factors underlying variation in their architecture. In grapevines, cluster density is an important trait that is used to assess grapevine crop quality and to forecast yield, in part because of the association between bunch density and fungal infestations such as Botrytis (Hed et al., 2009; Iland et al., 2011; Molitor & Beyer, 2014; Molitor et al., 2018). This study expands on previous work identifying variation in inflorescence architecture among cultivars (Shavrukov et al., 2004), finding notable differences in cluster architecture among species. A logical next step may be to use 3D images and PH with topological modeling to trace the development of inflorescences across multiple growing seasons in a mapping population. Methods presented here are also amenable to scanning with berries, provided some noteworthy technical challenges are first addressed (e.g. minimizing berry damage and rotating during transportation, cluster stabilization during scanning, and segmentation of 3D volumes with features that vary widely in their X-ray absorbance). This work would provide a more complete representation of cluster structure, as well as inform our berry potential simulation with genotype-specific empirical data. We plan to develop predictive structural models of grapevine cluster development using these techniques.

Imaging and shape analysis approaches presented here can also be used to tease apart subtle environmental influences on inflorescence architecture, and the major agronomic trait of bunch density. Identifying environmental effects on phenotypic variation has important implications both for vineyard management and the assessment of intra-clone variation across geographic space. Cluster compactness can be manipulated through a variety of agronomic practices (Molitor et al., 2012; Gil et al., 2013; Frioni et al., 2017; Gourieroux et al., 2017; Poni et al., 2018; Reeve et al., 2018). Techniques described here can be used to quantify influences of specific treatments on cluster architecture. In addition, because grapevines are clonally propagated, clusters from the same widespread clones can be collected from different geographic locations, scanned, and analysed for variation. High resolution assessment of inflorescence architecture offers important insights into natural variation in bunch density and the genetic and environmental factors that influence it. The capacity to capture 3D variation in this complex trait over space and time represents a promising advance for a valuable potential target of selection in one of the most economically important berry crops in the world.

Supplementary data

Supplementary data are available at JXB online.

Fig. S1. A maximum likelihood phylogenetic tree for 10 Vitis species.

Fig. S2. Summary of inflorescence geometric and topological traits and the distribution for 10 Vitis species.

Fig. S3. Comparison between X-ray scanned and simulated berry features for five commercially available table grape varieties.

Fig. S4. A neighbor joining phylogenetic tree for a subset of the Vitis dataset.

Fig. S5.Variation for each clade.

Fig. S6. Pairwise correlations of morphological traits (allo-metric relationships) showing linear regression lines for each species.

Fig. S7. Pairwise species classification.

Table S1. Trait description and calculation.

Table S2. Trait variance for each species.

Table S3. Trait loadings for two species classification.

Table S4. Trait Pagel’s lambda for phylogenetic analysis.

Table S5. Trait variation for each clade.

Video S1. Illustration of quantifying branching topology using persistent homology.

Video S2. Berry potential simulation.

Acknowledgements

The authors would like to acknowledge Elizabeth A. Kellogg (DDPSC) for valuable comments, particularly on phylogenetic analysis and inflorescence anatomy. We thank Noah Fahlgren (DDPSC) for computational assistance and Kari Miller (Washington University) for scanning assistance. We thank Zoë Migicovsky (Dalhousie University) for valuable comments, particularly on phylogenetic analysis and inflorescence anatomy.
comments. This work was supported by funding from the United States National Science Foundation projects IIA–1355406 and IOS–1638507 to CNT, and DBI–1759796 to AJM.

Author contributions

CNT, DHC, and JL designed the research; JL collected the samples and consulted on the biology; KD generated the X-ray data; LLK and AJM provided phylogenetic data and consulted for the biology; NJ and ML extracted pedicel diameter and angle; ML developed and extracted all the traits and conducted all the analysis and figures; ML, LLK, KD, JL, AJM, and CNT wrote the manuscript.

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